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PASSWORD:

TERMINAL (ENTER 1, 2, 3, OR ?):2

		,		-, -,,
* * *	* *	* *	* *	* Welcome to STN International * * * * * * * * *
NEWS	1			Web Page for STN Seminar Schedule - N. America
NEWS		AUG	06	CAS REGISTRY enhanced with new experimental property tags
NEWS	3	AUG		FSTA enhanced with new thesaurus edition
NEWS	4	AUG	13	CA/CAplus enhanced with additional kind codes for granted patents
NEWS	5	AUG	20	CA/CAplus enhanced with CAS indexing in pre-1907 records
NEWS		AUG		Full-text patent databases enhanced with predefined patent family display formats from INPADOCDB
NEWS	7	AUG	27	USPATOLD now available on STN
NEWS	8	AUG		CAS REGISTRY enhanced with additional experimental
MEMO	0	AUG	20	spectral property data
NEWS	9	SEP	07	STN AnaVist, Version 2.0, now available with Derwent
				World Patents Index
NEWS		SEP		FORIS renamed to SOFIS
NEWS		SEP		INPADOCDB enhanced with monthly SDI frequency
NEWS			17	1967–1998
NEWS	13	SEP	17	CAplus coverage extended to include traditional medicine patents
NEWS	1.4	SEP	24	EMBASE, EMBAL, and LEMBASE reloaded with enhancements
NEWS		OCT		CA/CAplus enhanced with pre-1907 records from Chemisches Zentralblatt
NEWS	16	OCT	10	BEILSTEIN updated with new compounds
NEWS		NOV		Derwent Indian patent publication number format enhanced
NEWS		NOV		WPIX enhanced with XML display format
NEWS		NOV		ICSD reloaded with enhancements
NEWS		DEC		LINPADOCDB now available on STN
NEWS				BEILSTEIN pricing structure to change
NEWS				USPATOLD added to additional database clusters
NEWS				IMSDRUGCONF removed from database clusters and STN
NEWS		DEC		DGENE now includes more than 10 million sequences
NEWS		DEC		TOXCENTER enhanced with 2008 MeSH vocabulary in MEDLINE segment
NEWS	20	DEC	17	MEDLINE segment MEDLINE and LMEDLINE updated with 2008 MeSH vocabulary
NEWS		DEC		CA/CAplus enhanced with new custom IPC display formats
		DEC		STN Viewer enhanced with full-text patent content
NEWS			-	from USPATOLD
NEWS		JAN		STN pricing information for 2008 now available
NEWS	30	JAN	16	CAS patent coverage enhanced to include exemplified prophetic substances
NEWS	31	JAN	28	USPATFULL, USPAT2, and USPATOLD enhanced with new custom IPC display formats
NEWS	32	JAN	28	MARPAT searching enhanced
NEWS	33	JAN	28	USGENE now provides USPTO sequence data within 3 days

of publication

NEWS 34 JAN 28 TOXCENTER enhanced with reloaded MEDLINE segment

NEWS 35 JAN 28 MEDLINE and LMEDLINE reloaded with enhancements

NEWS EXPRESS 19 SEPTEMBER 2007: CURRENT WINDOWS VERSION IS V8.2, CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),

AND CURRENT DISCOVER FILE IS DATED 19 SEPTEMBER 2007.

NEWS HOURS STN Operating Hours Plus Help Desk Availability

NEWS LOGIN Welcome Banner and News Items

NEWS IPC8 For general information regarding STN implementation of IPC 8

Enter NEWS followed by the item number or name to see news on that specific topic.

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* * * * * * * * * * * * * * * * STN Columbus * * * * * * * * * * * * * * * * * *

FILE 'HOME' ENTERED AT 09:04:35 ON 31 JAN 2008

=> fil req

COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION

FULL ESTIMATED COST 0.21 0.21

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STRUCTURE FILE UPDATES: 30 JAN 2008 HIGHEST RN 1001156-45-1 DICTIONARY FILE UPDATES: 30 JAN 2008 HIGHEST RN 1001156-45-1

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TSCA INFORMATION NOW CURRENT THROUGH June 29, 2007

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http://www.cas.org/support/stngen/stndoc/properties.html

=> e citric acid

E1 6 CITRIARBUSTI/BI 411 CITRIC/BI

E2 5411

E3 0 --> CITRIC ACID/BI F.4 18 CITRICA/BI

```
STN Search - 10/517,692
```

```
E5
              1 CITRICAL/BI
           5
1
                     CITRICARPA/BI
E6
E7
                     CITRICID/BI
E8
           4309
                     CITRICIDA/BI
E9
                     CITRICIDAL/BI
           1
E10
            1 7
                     CITRICIN/BI
E11
                     CITRICOCCUS/BI
         113
E12
                     CITRICOL/BI
=> e citric acid/cn
        1 CITRIC A-CYCLOHEXYLAMIDE/CN
E2
               1
                      CITRIC B-CYCLOHEXYLAMIDE/CN
E3
               1 --> CITRIC ACID/CN
E4
              1
                     CITRIC ACID 2-(TERT-BUTYL) 1,3-BIS(SUCCINIMIDYL) ESTER/CN
E5
                     CITRIC ACID 2-(TERT-BUTYL) ESTER/CN
              1
E6
               1
                     CITRIC ACID 2-METHYLIMIDAZOLE SALT/CN
E7
               1
                     CITRIC ACID 2-STEARYLOXYETHYL ESTER/CN
E8
               1
                     CITRIC ACID CALCIUM MAGNESIUM SALT/CN
               1
                     CITRIC ACID CHLORALIDE/CN
E9
                   CITRIC ACID CHLORALIDE/CN
CITRIC ACID DIAMIDE/CN
CITRIC ACID DIHYDRATE/CN
              1
E10
               1
E11
E12
               1
=> e malic acid/cn
E1 1 MALIBATOL A/CN
E2
                      MALIBATOL B/CN
E3
               1 --> MALIC ACID/CN
             1 --> MALIC ACID/CN

MALIC ACID 1-METHYL ESTER/CN

MALIC ACID 2-METHYLIMIDAZOLE SALT/CN

MALIC ACID CACTATE DICALORIDE/CN

MALIC ACID BARIUM SALT (1:1)/CN

MALIC ACID CHORALIDE/CN

MALIC ACID CHORALIDE/CN

MALIC ACID DEBYPORGENASE/CN

MALIC ACID DIBENZYL ESTER/CN

MALIC ACID DIBENZYL ESTER/CN

MALIC ACID DIETHANOLAMINE SALT/CN
E4
E5
E6
E7
E8
E9
E10
E11
E12
=> s e3
               1 "MALIC ACID"/CN
L1
=> e citric acid/cn
E1
       1
                     CITRIC A-CYCLOHEXYLAMIDE/CN
E2
               1
                      CITRIC B-CYCLOHEXYLAMIDE/CN
E3
               1 --> CITRIC ACID/CN
E4
                    CITRIC ACID 2-(TERT-BUTYL) 1,3-BIS(SUCCINIMIDYL) ESTER/CN
               1
                     CITRIC ACID 2-(TERT-BUTYL) ESTER/CN
E5
               1
                     CITRIC ACID 2-METHYLIMIDAZOLE SALT/CN
E6
               1
E7
               1
                     CITRIC ACID 2-STEARYLOXYETHYL ESTER/CN
               1
                     CITRIC ACID CALCIUM MAGNESIUM SALT/CN
E8
             1 CITRIC ACID CHLORALIDE/CN
1 CITRIC ACID CHLORALIDE/CN
1 CITRIC ACID DIAMIDE/CN
1 CITRIC ACID DIAMIDE/CN
E9
E11
E12
=> s e 3
         738466 E
       19827953 3
          15758 E 3
                    (E(W)3)
```

```
=> e oxalacetic acid/cn
           1 OXALACETATE-ASPARTATE AMINOTRANSFERASE/CN
E1
E2
                  OXALACETIC B-DECARBOXYLASE/CN
E3
            1 --> OXALACETIC ACID/CN
            1 OXALACETIC ACID 2-STILBAZOLE-4'-HYDRAZONE/CN
E4
                 OXALACETIC ACID DECARBOXYLASE/CN
E6
                 OXALACETIC ACID DIETHYL ESTER SODIUM SALT/CN
            OXALACETIC ACID O-METHYLOXIME/CN
OXALACETIC ACID RADICAL CATION/CN
OXALACETIC ACID, ((1-METHYL-3-OXO-1-BUTENYLAMINO) METHYLENE) -
E7
E8
E9
                  /CN
E10
            1
                  OXALACETIC ACID, ((1-METHYL-3-OXO-1-BUTENYLAMINO)METHYLENE)-
                   , DIETHYL ESTER/CN
                   OXALACETIC ACID, ((11B, 17-DIHYDROXY-3-OXOESTR-5(10)-EN-
E11
             1
                   17A-YL)METHYL) -, F-LACTONE, METHYL ESTER, CYCLIC
                    3-(ETHYLENE ACETAL)/CN
E12
             1
                   OXALACETIC ACID, ((11B, 17-DIHYDROXY-3-OXOESTR-5-EN-17.A
                   LPHA.-YL) METHYL) -, F-LACTONE, METHYL ESTER, CYCLIC 3-(
                   ETHYLENE ACETAL)/CN
=> s e3
             1 "OXALACETIC ACID"/CN
=> e citric acid/cn
            1
                  CITRIC A-CYCLOHEXYLAMIDE/CN
            1
E2
                  CITRIC B-CYCLOHEXYLAMIDE/CN
E3
            1 --> CITRIC ACID/CN
            1 CITRIC ACID 2-(TERT-BUTYL) 1,3-BIS(SUCCINIMIDYL) ESTER/CN
E4
E5
            1 CITRIC ACID 2-(TERT-BUTYL) ESTER/CN
E6
                 CITRIC ACID 2-METHYLIMIDAZOLE SALT/CN
            1
E7
                 CITRIC ACID 2-STEARYLOXYETHYL ESTER/CN
            1
E8
           1 CITRIC ACID CALCIUM MAGNESIUM SALT/CN
1 CITRIC ACID CHLORALIDE/CN
E9
E10
           1
                 CITRIC ACID CHLORIDE/CN
E11
            1
                 CITRIC ACID DIAMIDE/CN
E12
            1
                 CITRIC ACID DIHYDRATE/CN
=> s e3
             1 "CITRIC ACID"/CN
L4
=> e aconitic acid/cn
E1
             1
                  ACONITE, TINCTURE/CN
E2
             1
                  ACONITI TINCTURE/CN
E3
             1 --> ACONITIC ACID/CN
E4
                  ACONITIC ACID ANHYDRIDE-ETHYLENE-OCTYL ACRYLATE-PROPYLENE GR
             1
                  AFT COPOLYMER/CN
E5
                  ACONITIC ACID IRON SALT/CN
             1
                  ACONITIC ACID MONOMETHYL ESTER/CN
E6
             1
                  ACONITIC ACID TRIBENZYL ESTER/CN
E7
             1
                  ACONITIC ACID, A-AMINO-, TRIETHYL ESTER/CN
E8
             1
                  ACONITIC ACID, A-BROMO-, TRIETHYL ESTER/CN
E9
             1
E10
            1
                  ACONITIC ACID, A-CYANO-Γ-FLUORO-, TRIETHYL ESTER
E11
            1
                   ACONITIC ACID, A-ETHOXY-F-OXO-, TRIETHYL ESTER/C
E12
             1
                  ACONITIC ACID, A-PROPOXY-, TRIETHYL ESTER/CN
=> s e3
           1 "ACONITIC ACID"/CN
1.5
```

```
=> e malate
E1
            1
                  MALATASE/BI
MALATATE/BI
E2
            1
E3
          5352 --> MALATE/BI
            1 MALATE63/BI
E4
E5
            1
                  MALATE:NA+/BI
           4 MALATE:NAD+/BI
87 MALATE:QUINONE/BI
1 MALATEDEHYDROGEN/BI
1 MALATEDEHYDROGENASE/BI
E6
E7
E8
E9
E10
            4
                  MALATES/BI
                  MALATESTAI/BI
E11
            1
E12
            2
                  MALATESYN/BI
=> s e3
          5352 MALATE/BI
L6
=> d his
     (FILE 'HOME' ENTERED AT 09:04:35 ON 31 JAN 2008)
     FILE 'REGISTRY' ENTERED AT 09:04:41 ON 31 JAN 2008
                E CITRIC ACID
                E CITRIC ACID/CN
                E MALIC ACID/CN
               1 S E3
                E CITRIC ACID/CN
          15758 S E 3
                E OXALACETIC ACID/CN
               1 S E3
                E CITRIC ACID/CN
              1 S E3
T. 4
                E ACONITIC ACID/CN
1.5
               1 S E3
                E MALATE
L6
           5352 S E3
=> fil caplus
COST IN U.S. DOLLARS
                                                   SINCE FILE
                                                                    TOTAL.
                                                         ENTRY
                                                                 SESSION
FULL ESTIMATED COST
                                                         39.73
                                                                    39.94
FILE 'CAPLUS' ENTERED AT 09:09:21 ON 31 JAN 2008
```

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```
FILE COVERS 1907 - 31 Jan 2008 VOL 148 ISS 5
FILE LAST UPDATED: 30 Jan 2008 (20080130/ED)
```

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=> \$ 11 or 12 or 13 or 14 or 15 or 16
22764 L1
920942 L2
4146 L3
68175 L4
1003 L5
22725 L6
L7 1016908 L1 OR L2 OR L3 OR L4 OR L5 OR L6

=> s 11 L8 22764 L1

=> s 12 L9 920942 L2

=> s 13 L10 4146 L3

=> s 14 L11 68175 L4

=> s 15 L12 1003 L5

=> s 16 L13

=> fil reg COST IN U.S. DOLLARS

22725 L6

SINCE FILE ENTRY 0.48

TOTAL SESSION 40.42

FULL ESTIMATED COST

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http://www.cas.org/support/stngen/stndoc/properties.html

```
=> e hydroxysuccinimide
E1
           2
                 HYDROXYSUCCINIMI/BI
E2
           49
                  HYDROXYSUCCINIMID/BI
E3
           222 --> HYDROXYSUCCINIMIDE/BI
E4
                 HYDROXYSUCCINIMIDESTER/BI
E5
           17
                 HYDROXYSUCCINIMIDO/BI
E6
           1
                 HYDROXYSUCCINIMIDOTHALLIUM/BI
E7
            4
                 HYDROXYSUCCINIMIDOYL/BI
E8
           49
                 HYDROXYSUCCINIMIDYL/BI
E9
            2
                 HYDROXYSUCCINIMIDYLPROPION/BI
E10
            2
                 HYDROXYSUCCINIMIDYLPROPIONATE/BI
E11
                 HYDROXYSUCCINIMIDYLUNDECAN/BI
            1
            1
E12
                 HYDROXYSUCCINIMIDYLUNDECANO/BI
=> e n-hydroxysuccinimide/cn
                  N-HYDROXYSUCCINAMIC ACID/CN
E2
                  N-HYDROXYSUCCINAMIDE/CN
E3
             1 --> N-HYDROXYSUCCINIMIDE/CN
E4
                  N-HYDROXYSUCCINIMIDE 4-AZIDO-2-HYDROXYBENZOATE/CN
E5
                  N-HYDROXYSUCCINIMIDE 4-AZIDOBENZOATE/CN
                 N-HYDROXYSUCCINIMIDE 4-AZIDOBENZOIC ESTER/CN
E6
                 N-HYDROXYSUCCINIMIDE ACETATE/CN
                 N-HYDROXYSUCCINIMIDE BROMOACETATE/CN
E8
                 N-HYDROXYSUCCINIMIDE CHLOROFORMATE/CN
E9
                 N-HYDROXYSUCCINIMIDE DOCOSANOATE/CN
E10
                 N-HYDROXYSUCCINIMIDE ESTER OF 2-NITRO-5-AZIDOBENZOYL-GLYCINE
E11
            1
                  N-HYDROXYSUCCINIMIDE ESTER OF N-(4-CARBOXYPHENYLMETHYL) MALEI
                  MIDE/CN
=> s e3
L14
             1 N-HYDROXYSUCCINIMIDE/CN
=> e n-hydroxysulfosuccinimide/cn
E1
                  N-HYDROXYSUCCINIMIDYL PYRENEBUTANOATE/CN
            1
E2
                  N-HYDROXYSULFONAPHTHALIMIDE/CN
E3
             1 --> N-HYDROXYSULFOSUCCINIMIDE/CN
E4
                 N-HYDROXYSULFOSUCCINIMIDE SODIUM SALT/CN
E5
                 N-HYDROXYSULFOSUCCINIMIDYL-DOTA/CN
E6
                 N-HYDROXYTETRABROMOPHTHALIMIDE/CN
E7
                 N-HYDROXYTETRACHLOROPHTHALIMIDE/CN
E8
                 N-HYDROXYTETRADECANAMIDE/CN
E9
                 N-HYDROXYTETRAPROPENYLSUCCINIMIDE/CN
                 N-HYDROXYTHIAZOLE-2(3H)-THIONE/CN
E10
E11
                 N-HYDROXYTHIOBENZANILIDE/CN
                 N-HYDROXYTHIOCARBANILIDE/CN
=> s e3
L15
             1 N-HYDROXYSULFOSUCCINIMIDE/CN
=> fil caplus
COST IN U.S. DOLLARS
                                                SINCE FILE
                                                                TOTAL
                                                     ENTRY
                                                             SESSION
FULL ESTIMATED COST
                                                     10.76
                                                               51.18
FILE 'CAPLUS' ENTERED AT 09:10:58 ON 31 JAN 2008
```

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```
FILE COVERS 1907 - 31 Jan 2008 VOL 148 ISS 5
FILE LAST UPDATED: 30 Jan 2008 (20080130/ED)
```

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http://www.cas.org/infopolicy.html

5280 L14

=> s 114

L16

```
=> s 115
L17
          312 L15
=> s 116 or 117
         5501 L16 OR L17
L18
=> s 115 and (pv<=2003)
           312 L15
      23975525 PY<=2003
L19
          162 L15 AND (PY<=2003)
=> d his
     (FILE 'HOME' ENTERED AT 09:04:35 ON 31 JAN 2008)
     FILE 'REGISTRY' ENTERED AT 09:04:41 ON 31 JAN 2008
                E CITRIC ACID
                E CITRIC ACID/CN
                E MALIC ACID/CN
              1 S E3
                E CITRIC ACID/CN
          15758 S E 3
                E OXALACETIC ACID/CN
              1 S E3
                E CITRIC ACID/CN
L4
              1 S E3
                E ACONITIC ACID/CN
L5
              1 S E3
                E MALATE
1.6
           5352 S E3
    FILE 'CAPLUS' ENTERED AT 09:09:21 ON 31 JAN 2008
       1016908 S L1 OR L2 OR L3 OR L4 OR L5 OR L6
T.R
         22764 S L1
L9
        920942 S L2
```

```
STN Search - 10/517,692
L10
          4146 S L3
1,11
         68175 S L4
L12
          1003 S L5
L13
          22725 S L6
     FILE 'REGISTRY' ENTERED AT 09:10:08 ON 31 JAN 2008
                E HYDROXYSUCCINIMIDE
                E N-HYDROXYSUCCINIMIDE/CN
L14
              1 S E3
               E N-HYDROXYSULFOSUCCINIMIDE/CN
L15
              1 S E3
     FILE 'CAPLUS' ENTERED AT 09:10:58 ON 31 JAN 2008
1.16
          5280 S L14
L17
           312 S L15
L18
          5501 S L16 OR L17
L19
           162 S L15 AND (PY<=2003)
=> s 17 and (pv<=2003)
      23975525 PY<=2003
L20
        784186 L7 AND (PY<=2003)
=> s 18 and (py<=2003)
      23975525 PY<=2003
        18251 L8 AND (PY<=2003)
=> s 19 and (py<=2003)
      23975525 PY<=2003
       707903 L9 AND (PY<=2003)
=> s 110 and (py<=2003)
      23975525 PY<=2003
L23
         3763 L10 AND (PY<=2003)
=> s 111 and (py<=2003)
      23975525 PY<=2003
L24
         50287 L11 AND (PY<=2003)
=> s 112 and (pv<=2003)
      23975525 PY<=2003
          890 L12 AND (PY<=2003)
=> s 113 and (py<=2003)
      23975525 PY<=2003
1.26
        19656 L13 AND (PY<=2003)
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     (FILE 'HOME' ENTERED AT 09:04:35 ON 31 JAN 2008)
     FILE 'REGISTRY' ENTERED AT 09:04:41 ON 31 JAN 2008
                E CITRIC ACID
                E CITRIC ACID/CN
                E MALIC ACID/CN
              1 S E3
                E CITRIC ACID/CN
1.2
          15758 S E 3
               E OXALACETIC ACID/CN
              1 S E3
                E CITRIC ACID/CN
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| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|---------------|------|----------|-----------------|------------|
| | | | | |
| WO 2003104814 | A2 | 20031218 | WO 2003-US14503 | 20030508 < |

PCT Int. Appl., 122 pp.

CODEN: PIXXD2

Patent

English

SOURCE:

LANGUAGE:

DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: 8 PATENT INFORMATION:

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WO 2003104814
                        A3 20041111
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
             PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ,
             UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
             KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES,
             FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
             BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                             20031222 AU 2003-233498 20030508 <--
20050330 EP 2003-728775 20030508
     AU 2003233498
                         A1
     EP 1518115
                         A2
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
                                            JP 2004-511834
    JP 2005529335
                        T 20050929
                                                                   20030508
                                                              P 20020610
                                            US 2002-388120P P 20021016
US 2002-419136P P 20021016
TE 2002-434061P P 20021217
                                            US 2002-388120P
PRIORITY APPLN. INFO.:
                                            US 2003-447605P
                                                                P 20030214
                                            WO 2003-US14503
                                                                W 20030508
    An open capillary channel device for open tubular solid phase extraction of
    mols. capable of providing a tube enrichment factor of at least 1. The
     device comprises a channel having one end connected to a pump for pumping
     liquid and gas, and the other end can be connected to an interface for a
     protein chip sample applicator or a mass spectrometer. The inner surface
     of the channel, an extraction surface, can be bonded to an affinity binding
     agent such as a chelated metal, a protein, a sugar or nucleic acid. The
    method uses this device to bind analyte mols. from a sample solution to the
    affinity extraction surface and desorb analyte from the extraction surface
```

with a desorbent liquid, with an extraction factor greater than 1.

L28 ANSWER 2 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

```
ACCESSION NUMBER:
                      2003:118290 CAPLUS
DOCUMENT NUMBER:
                       138:177983
TITLE:
                       Upconversion luminescence materials and methods of
                       making and using same
INVENTOR(S):
                       Chen, Wei
PATENT ASSIGNEE(S):
                      USA
SOURCE:
                       U.S. Pat. Appl. Publ., 40 pp.
                       CODEN: USXXCO
DOCUMENT TYPE:
                       Patent
LANGUAGE:
                       English
```

FAMILY ACC. NUM. COUNT: 3 PATENT INFORMATION:

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------|--------------------|------|----------|-------------------|------------|
| | | | | | |
| | US 2003030067 | A1 | 20030213 | US 2002-166313 | 20020606 < |
| | US 7008559 | B2 | 20060307 | | |
| | US 2003064532 | A1 | 20030403 | US 2002-223764 | 20020819 < |
| | US 7067072 | B2 | 20060627 | | |
| | US 2005169348 | A1 | 20050804 | US 2003-460531 | 20030612 |
| | US 2006274813 | A9 | 20061207 | | |
| | US 2005253095 | A1 | 20051117 | US 2005-67373 | 20050225 |
| | US 7126136 | B2 | 20061024 | | |
| | US 2006140240 | A1 | 20060629 | US 2005-202005 | 20050811 |
| PRIO | RITY APPLN. INFO.: | | | US 2001-296333P P | 20010606 |
| | | | | US 2002-356598P P | 20020211 |
| | | | | | |

| US | 2001-313236P | P | 20010817 |
|----|--------------|----|----------|
| US | 2002-356542P | P | 20020211 |
| US | 2002-166313 | A2 | 20020606 |
| US | 2002-388211P | P | 20020612 |
| US | 2002-223764 | A1 | 20020819 |

AB An upconversion luminescence material of the general formula X:Y (X:host; Y:dopant) wherein the at least one dopant is capable of increasing the luminescence intensity or quantum efficiency of the host is described wherein X may be a semiconductor nanoparticle selected from ZnSx, ZnSex, ZnTex, CdSx, CdSex, CdTex, PbSx, PbSex, PbTex, MqSx, CaSx, BaSx, SrSx and Y may be selected from Eu3+, Tb3+, Ce3+, Er3+, Mn2+ and Cu+. An upconversion luminescence production assembly is also described comprising an electromagnetic source emitting an excitation having an excitation wavelength; a substrate positioned within the excitation emitted by the electromagnetic source; and a upconversion luminescent (UCL) material operably associated with at least a portion of the substrate such that the excitation emitted by the electromagnetic source is received by at least a portion of the UCL material, the UCL material producing an emission through upconversion luminescence having an emission wavelength shorter than the excitation wavelength of the excitation received by the UCL material. Use of the phosphor in biol, and biomedical devices is indicated.

REFERENCE COUNT: 250 THERE ARE 250 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

L28 ANSWER 3 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2003:43871 CAPLUS

DOCUMENT NUMBER: 138:364590

TITLE: Kinetic locking-on and auxiliary tactics for bioaffinity purification of NADP+-dependent dehydrogenases using N6-linked immobilized NADP+ derivatives: studies with mammalian and microbial

glutamate dehydrogenases

AUTHOR(S): McMahon, Mary; Tynan, Julie; Mulcahy, Patricia

CORPORATE SOURCE: Department of Applied Biology and Chemistry, Institute of Technology, Carlow, Ire.

Biotechnology and Bioengineering (2003), 81(3),

356-369

CODEN: BIBIAU: ISSN: 0006-3592

PUBLISHER: John Wiley & Sons, Inc.

DOCUMENT TYPE: Journal

SOURCE:

AB

LANGUAGE:

English This study is concerned with the development and application of kinetic locking-on and auxiliary tactics for bioaffinity purification of NADP+-dependent dehydrogenases, specifically (1) the synthesis and characterization of highly substituted N6-linked immobilized NADP+ derivs. using a rapid solid-phase modular approach; (2) the evaluation of the N6-linked immobilized NADP+ derivs. for use with the kinetic locking-on strategy for bioaffinity purification of NADP+-dependent dehydrogenases: Model bioaffinity chromatog, studies with glutamate dehydrogenase from bovine liver (GDH with dual cofactor specificity, EC 1.4.1.3) and glutamate dehydrogenase from Candida utilis (GDH which is NADP+-specific, EC 1.4.1.4); (3) the selection of an effective "stripping ligand" for NADP+-dehydrogenase bioaffinity purifications using N6-linked immobilized NADP+ derivs. in the locking-on mode; and (4) the application of the developed bioaffinity chromatog. system to the purification of C. utilis GDH from a crude cellular extract Results confirm that the newly developed N6-linked immobilized NADP+ derivs. are suitable for the one-step bioaffinity purification of NADP+-dependent GDH provided that they are used in

the locking-on mode, steps are taken to inhibit alkaline phosphatase activity in crude cellular exts., and 2',5'-ADP is used as the stripping ligand during chromatog. The general principles described here are supported by a specific sample enzyme purification; the purification of C. utilis GDH to electrophoretic homogeneity in a single bioaffinity chromatog, step (specific activity, 9.12 μmol/min/mg; purification factor, 83.7; yield 88%). The potential for development of analogous bioaffinity systems for other NADP+-dependent dehydrogenases is also discussed.

REFERENCE COUNT: 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L28 ANSWER 4 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2002:825899 CAPLUS

DOCUMENT NUMBER: 138:113964

TITLE: Preparation, characterization and application of

alkanethiol self-assembled monolayers modified with tetrathiafulvalene and glucose oxidase at a gold disk

electrode

Campuzano, Susana; Galvez, Rocio; Pedrero, Maria; De AUTHOR(S): Villena, F. Javier Manuel; Pingarron, Jose M.

Doto, Ouimica Analitica, Facultad de CC, Ouimicas, CORPORATE SOURCE: Universidad Complutense de Madrid, Madrid, E-28040,

Spain

Proceedings - Electrochemical Society (2001), SOURCE:

2001-18(Chemical and Biological Sensors and Analytical

THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS

Methods II), 602-608

CODEN: PESODO; ISSN: 0161-6374

PUBLISHER: Electrochemical Society

Journal DOCUMENT TYPE: LANGUAGE: English

In this work, the results obtained with a gold disk electrode modified with alkanethiol self-assembled monolayers (SAMs), and glucose oxidase (GOD), and the redox mediator tetrathiafulvalene (TTF) immobilized atop are presented. Thus, a gold electrode modified with a mercaptopropionic acid SAM, where GOD and TTF were immobilized by crosslinking with glutaraldehyde, allowed linear calibration curves for glucose, obtained by amperometry in stirred solns. at an applied potential of +0.20 V, in the 5.0 10-6 - 1.0 10-2 mol L-1 range. A detection limit of 1.3 10-6 mol L-1, and a RSD of 5.2% (n=10), at a concentration level of 1.0 10-4 mol L-1, were found. No leaching of the enzyme and mediator is observed during the whole working day. The modified electrode is stable in dry conditions for 24 h

and for at least 100 h if kept in a 4 °C H2PO4-/HPO42- buffer solution (pH 7.4).

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

8 L28 ANSWER 5 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2001:763341 CAPLUS

DOCUMENT NUMBER: 135:312579

TITLE: Magnetically-responsive microspheres INVENTOR(S): Chandler, Donald J.; Herren, Michael A.

Luminex Corporation, USA PATENT ASSIGNEE(S): SOURCE: PCT Int. Appl., 16 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

REFERENCE COUNT:

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WO 2001078087 A2 20011018 WO 2001-US11122 WO 2001078087 A3 20020704
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             HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
             LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
             SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,
             YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
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             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
     US 2001046602 A1 20011129 US 2001-826960
     US 6773812
                         B2 20040810
PRIORITY APPLN. INFO.:
                                           US 2000-194889P
                                                              P 20000406
AB Microspheres are constructed using magnetic particles. Hybrid
    microspheres are constructed using fluorescent or luminescent microspheres
     and magnetic nanoparticles. Reactive moieties on the surface of the
     resultant particles can be used for attachment of biol. active mols., thus
     allowing selective sepns, and anal, assays to be performed.
     Distinguishable subsets of microspheres can be constructed based on
     fluorescent intensities, and sepns. can be affected based on variable
     degree of magnetic content.
L28 ANSWER 6 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER: 2001:163317 CAPLUS
DOCUMENT NUMBER:
                        134:339824
TITLE:
                        A novel chitosan derivative to immobilize
                        α-L-rhamnopyranosidase from Aspergillus niger
                        for application in beverage technologies
                        Spagna, G.; Barbagallo, R. N.; Casarini, D.; Pifferi,
AUTHOR(S):
                        P. G.
CORPORATE SOURCE:
                        Food Biotechnology Group from the Department of
                        Horticulture, Floriculture, Arboriculture and
                        Agroindustrial Technology (DOFATA), University of
                        Catania, Catania, 95123, Italy
SOURCE:
                        Enzyme and Microbial Technology (2001), 28(4-5),
                        427-438
                        CODEN: EMTED2; ISSN: 0141-0229
PUBLISHER:
                        Elsevier Science Ireland Ltd.
DOCUMENT TYPE:
                        Journal
LANGUAGE:
                        English
AB
    α-L-rhamnopyranosidase (Rha, EC 3.2.1.40) is an enzyme of
    considerable importance to food technol. in increasing the aroma of wines,
     musts, fruit juices and other beverages. The aim of this research is the
     immobilization of the Rha contained in a com. preparation already used in the
     winemaking industry. The immobilization supports tested were chitin,
    chitosan and derivatized chitosan, diethylaminoethyl chitosan
    (DE-chitosan) never previously used for this type of application.
     Particularly, on DE-chitosan, the Rha was adsorbed and cross-linked with
     various bifunctional agents (glutaraldehyde, diepoxyoctane, suberimidate
     and carbodiimide), whose best results (immobilization yields and activity)
    were obtained with carbodiimide (EDC) that allowed a reduction in the
    involvement of the enzyme amine groups that are probably important in
    catalytic mechanism. In addition, the use of rhamnose and a succinimide
    (NHS) during crosslinking enhanced the action of the EDC and so increased
     the immobilization yield and activity. The immobilized Rha retained the
    kinetic parameters (Km and Vmax) of the free enzyme and increased
```

stability. Moreover, this biocatalyst allowed an increase in the aroma in a model wine solution containing glycosidic precursors with a marked reduction

specificity toward tertiary monoterpenols as compared to the free enzyme.

REFERENCE COUNT: 60 THERE ARE 60 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L28 ANSWER 7 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1995:978863 CAPLUS

DOCUMENT NUMBER: 124:3993

TITLE: Solid phase immunoassay to detect inhibitors of proteolytic enzymes using a tubulin substrate

INVENTOR(S): Islam, Khalid; Carrano, Lucia; Denaro, Maurizio
PATENT ASSIGNEE(S): Gruppo Lepetit S.p.A., Italy

SOURCE: PCT Int. Appl., 28 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

| | IT NO. | | KI | ND DATE | APPLICATION NO. | DATE |
|------------|-------------------|--------|--------|-------------|--|--------------|
| WO 95 | 26505 | | A: | 19951005 | WO 1995-EP867 | 19950309 < |
| | V: JP,
RW: AT. | | H. DE. | DK. ES. FR. | GB, GR, IE, IT, LU, MC | , NL. PT. SE |
| EP 75 | | , | A: | | EP 1995-913069 | |
| | 3152 | | В: | | | |
| | R: AT, | BE, C | H, DE, | | GB, GR, IE, IT, LI, LU
JP 1995-524925 | |
| | 9510786
517712 | | B: | | | 19950309 < |
| AT 16 | | | T. | | | 19950309 < |
| ES 21 | 114743 | | T | 19980601 | ES 1995-913069 | 19950309 < |
| | 159746 | | A | 20001212 | | 19960923 < |
| PRIORITY A | APPLN. | INFO.: | | | EP 1994-104922 | A 19940329 |
| | | | | | WO 1995-EP867 | W 19950309 |

AR A solid phase immunoassay for detecting specific inhibitors of proteolytic enzymes in biol. fluids or in any kind of solution containing them, as well as for detecting proteolytic activities in any solution containing them, is presented. The assay allows determination of inhibitors of the more common classes of proteases at the same time, using the same peptide substrate and the same detection antibody. Tubulin protein or a tubulin-like peptide covalently linked to a suitable support is contacted with a solution containing the proteolytic activity together with a protease inhibitor. Inhibitor activity against the selected proteases is determined by contacting the support with a solution containing a labeled monoclonal antibody which specifically recognizes the free end of the tubulin protein linked to the support. The method is illustrated using tubulin or a 21-residues containing the C-terminus of α-tubulin covalently linked to plastic microtiter wells via bis(sulfosuccinimidyl)suberate. The antibody preparation consists of rat antibody YL 1/2 specific for the C-terminus of undegraded, linked tubulin (or the synthetic peptide) and a peroxidase-labeled anti-YL 1/2 antibody. The method is accurate, precise, rapid, and easy to practice, and the intra- and inter- assay precision are well within the range of values currently accepted for anal. purposes.

L28 ANSWER 8 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1993:467303 CAPLUS

DOCUMENT NUMBER: 119:67303

TITLE: Reactive chitosan-coated articles and test kit for

immunoassay

INVENTOR(S): Saunders, Mary S.; Pegg, Randall K.

PATENT ASSIGNEE(S): US.

SOURCE: U.S., 5 pp. CODEN: USXXAM DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

KIND DATE APPLICATION NO. PATENT NO. -----A 19930504 US 1991-662420 19910228 <--US 1991-662420 19910228 US 5208166 PRIORITY APPLN. INFO.: AB A solid surface is coated with chitosan and a polyvalent organic acid, and the chitosan is oxidized to provide a substratum for immobilization of

immunochem. reagents for use in immunoassays. Thus, a stock solution of chitosan (0.02 g/mL in 0.1M citric acid, pH 2.0) was diluted 1:10, used to coat a polystyrene microtiter strip, and the chitosan was oxidized with NaNO2 (0.002 g/200 mL) for immobilization of rabbit IgG.

=> d his

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FILE 'REGISTRY' ENTERED AT 09:04:41 ON 31 JAN 2008

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E MALIC ACID/CN

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E CITRIC ACID/CN

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1 S E3 E CITRIC ACID/CN

T. 4 1 S E3 E ACONITIC ACID/CN L5 1 S E3

E MALATE 1.6 5352 S E3

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L12 1003 S L5 L13 22725 S L6

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L15 1 S E3

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QUALIFICATION NOT VALID FOR NUMERIC DATA 'PY/RACT' Numeric data cannot be field qualified.

=> s 17/ract

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You may have tried to apply a field code to a term that already has a field code. You can only add a field code to a term that has no field code appended to it.

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ENTRY
46.00 | TOTAL
SESSION
97.18 |
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| DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) | SINCE FILE
ENTRY | TOTAL
SESSION |
| CA SUBSCRIBER PRICE | -6.40 | -6.40 |

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=> e malic acid/prep

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file. To see a list of valid EXPAND field codes, enter HELP
SFIELDS at an arrow prompt (=>).

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E4

E5 E6

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1 MALIC ACID 2-METHYLIMIDAZOLE SALT/CN
1 MALIC ACID CECTATE DICHLORIDE/CN
1 MALIC ACID BARIUM SALT (1:1)/CN
1 MALIC ACID CHURALIDE/CN
1 MALIC ACID DEHYDROGENASE/CN E7 E8 E9

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ENTRY

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STN Search - 10/517,692
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                E MALIC ACID/CN
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               E CITRIC ACID/CN
          15758 S E 3
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               E OXALACETIC ACID/CN
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               E CITRIC ACID/CN
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                E N-HYDROXYSUCCINIMIDE/CN
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T.15

1 S E3

1 S E3

E N-HYDROXYSULFOSUCCINIMIDE/CN

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          3763 S L10 AND (PY<=2003)
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L31
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=> s 115/ract
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       3069601 RACT/RL
L37
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                 (L15 (L) RACT/RL)
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L38
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L41
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L42
          48 L40 AND L41
=> d ibib abs 1-48
L42 ANSWER 1 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                        2006:889280 CAPLUS
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DOCUMENT NUMBER: 145:299763

TITLE: Devices with multiple surface functionality coated

with phosphates or phosphonates

INVENTOR(S): Schwartz, Jeffrey; Gawalt, Ellen S.; Alvatroni,

Michael J.

PATENT ASSIGNEE(S): Princeton University, USA

SOURCE: U.S. Pat. Appl. Publ., 57pp., Cont.-in-part of U.S.

Ser. No. 876,294. CODEN: USXXCO

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 8
PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | | DATE |
|---|--|------|--|-----------------------------------|----------------------|
| PATENT NO. US 2006194008 US 6645644 US 2004001999 US 2004021999 US 2004265571 US 2005031910 PRIORITY APPLN. INFO.: | KIND

A1
B1
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A1
A1 | DATE | APPLICATION NO. US 2006-330814 US 2000-68080 US 2002-179743 US 2003-405557 US 2003-701591 US 2004-876298 US 2909-688080 US 2001-300144P US 2002-369236P US 2002-369236P US 2002-369237P US 2002-369237P US 2003-446680P US 2003-446681P US 2003-446681P US 2003-446681P US 2003-446681P US 2003-446681P US 2003-446681P US 2003-465318P | A2
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P | DATE |
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US 2003-490613P | P
P | 20030623
20030728 |
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US 2004-876294 | | 20031104 20040623 |
| | | | US 2005-643647P | P | 20050113 |
| | | | US 2005-643648P | P | 20050113 |
| | | | US 2005-684159P | P | 20050525 |
| | | | US 2005-699498P | P | 20050715 |
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| | | | US 1997-35040P | P | 19970113 |

plurality of phosphonate moieties, or both, covalently bonded to an oxide surface of an implantable substrate are provided. The coatings exhibit one or more of the following characteristics: (a) the surface phosphorus-containing group d. of the coated regions of the substrate is at least about 0.1 nmol/cm2; (b) the phosphorus-based coating has a thickness of less than about 10 nm; or (c) the surface phosphorus-containing group d. of the coated regions of the substrate is equal to or greater than the surface hydroxyl group d. of the oxide surface of the substrate. Implantable devices embodying the coated substrates are also disclosed. Thus, regions of a titanium hip implant were coated with (1) 11-hydroxyundecylphosphonic acid to which an osteoconductive mol. such as a peptide containing the RGD moiety is attached, to induce osteoconduction,

AB Phosphorus-based coatings having a plurality of phosphate moieties, a

US 1997-794833

A2 19970204

a peptide containing the RGD moiety is attached, to induce osteoconduction (2) underivatized 11-hydroxyundecylphosphonic acid, to prevent corrosion and leaching of metals, and (3) octadecylphosphonic acid, to lubricate the

interface between the ball and interior surface of the acetabular cup and to minimize wear debris generated from abrasion at the interface between the surfaces.

L42 ANSWER 2 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2004:501609 CAPLUS

DOCUMENT NUMBER: 142:172650

TITLE: Labelling technique of biomolecules for target

radiotherapy

Bai, Hongsheng; Jin, Xiaohai; Zhen, Cheng; Jia, Bing AUTHOR(S):

Fan Honggiang; Lu, Weiwei

CORPORATE SOURCE: Department of Isotope, China Institute of Atomic

Energy, Beijing, Peop. Rep. China

International Atomic Energy Agency, [Technical SOURCE: Document], IAEA-TECDOC (2003), IAEA-TECDOC-1359,

Labeling Techniques of Biomolecules for Targeted

Radiotherapy, 65-71

CODEN: IAEIE2; ISSN: 1011-4289

DOCUMENT TYPE: Report LANGUAGE: English

Labeling techniques were developed for the preparation of biomols. (DOTA-IgG, DOTA-lanreotide, anti-hepatoma antibody fragment, lanreotide) with

radionuclides such as 90Y, 153Sm and 188Re. The labeling yield and radiochem. purity of these labeling biomols. were determined by PC, ITLC and

Sep-Pak C18 cartridge. The stability in vitro and bio-behavior in normal rats were also evaluated. The exptl. results showed that labeling efficiency of biomols. (DOTA-IgG and DOTA-lanreotide) with 90Y and 153Sm

is more than 95% and had good stability in vitro, but the labeling efficiency of biomols. (anti-hepatoma antibody fragment and lanreotide) with 188Re via directly labeling technique is at range of 88% .apprx. 95%

and stability in vitro was less.

L42 ANSWER 3 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2003:918360 CAPLUS

DOCUMENT NUMBER: 140:281056

TITLE: Vasorelaxant activity of N-caffeoylamino acids AUTHOR(S): Iizuka, Toru; Funayama, Hiroko; Kusano, Genjiro;

Nagai, Masahiro

Fac. of Pharmaceutical Sciences, Hoshi Univ., Tokyo, CORPORATE SOURCE:

142-8501, Japan

SOURCE: Yakugaku Zasshi (2003), 123(11), 963-971

> CODEN: YKKZAJ; ISSN: 0031-6903 Pharmaceutical Society of Japan

PUBLISHER: DOCUMENT TYPE: Journal

LANGUAGE: Japanese

Twelve N-caffeovlamino acids and N-cinnamovlamino acids were synthesized and their vasorelaxation activity against norepinephrine (NE)-induced contraction of rat aorta was examined The following structure-activity relationships were found. (1) On the benzene ring, the caffeoyl structure is effective for vasorelaxation, while the cinnamoyl structure reduced vasorelaxation activity. (2) Four to six carbons are more effective as the carbon chain connecting the acylamino and carboxyl terminal groups. N-Caffeoyl- β -alanine and N-caffeoyltranexamic acid were used to investigate the action mechanism of vasorelaxing activities. It is believed that these compds. antagonize NE-induced vasocontraction by inhibiting receptor-operated calcium channels.

L42 ANSWER 4 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2003:891635 CAPLUS

DOCUMENT NUMBER: 140:402703 TITLE: Immobilized culture of nonadherent cells on an olevl

poly(ethylene glycol) ether-modified surface Kato, Koichi; Umezawa, Kohei; Funeriu, Daniel P.; AUTHOR(S):

Miyake, Masato; Miyake, Jun; Nagamune, Teruyuki

National Institute of Advanced Industrial Science and CORPORATE SOURCE:

Technology, Hyogo, Japan

BioTechniques (2003), 35(5), 1014-1016,1018,1020-1021 SOURCE:

CODEN: BINODO: ISSN: 0736-6205

PUBLISHER: Eaton Publishing Co.

DOCUMENT TYPE: Journal

LANGUAGE: English

Microarrays of living cells are an emerging tool in systems such as reverse transfection. These studies are limited to adherent cells partly because of the difficulty of cell immobilization. Using a newly developed reagent, the biocompatible anchor for membrane (BAM), the rapid and strong attachment of living nonadherent cells and adherent cells on BAM-modified

surfaces is shown in the study. Normal cellular growth was observed for over 7 days on BAM-modified surfaces. It is expected that this methodol. to

greatly expand the scope of current cell microarray technol.

REFERENCE COUNT: THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L42 ANSWER 5 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2003:707991 CAPLUS DOCUMENT NUMBER: 140:4873

TITLE: Self-condensation of activated malonic acid half esters: a model for the decarboxylative Claisen

condensation in polyketide biosynthesis

AUTHOR(S): Ryu, Youngha; Scott, A. Ian

CORPORATE SOURCE: Department of Chemistry, Center for Biological NMR,

Texas A&M University,

College Station, TX, 77843, USA

SOURCE: Tetrahedron Letters (2003), 44(40), 7499-7502

CODEN: TELEAY: ISSN: 0040-4039

PUBLISHER: Elsevier Science B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 140:4873

The reaction of a malonic acid half oxyesters RO2CCH2CO2H [R = CH2Ph, Ph,

(E)-CH2CH:CMe(CH2)2CH:CMe2, etc.] with a N-hydroxysuccinimidyl ester-forming reagent (O-(N-succinimidyl)-N,N,N',N'-tetramethyluronium

tetrafluoroborate) resulted in self-condensation to provide the

corresponding 1,3-acetonedicarboxylic acid diesters RO2CCH2COCH2CO2R. This new method does not require a divalent metal chelator or a

coordinating solvent for successful condensation.

REFERENCE COUNT: THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS 14 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L42 ANSWER 6 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2002:971937 CAPLUS DOCUMENT NUMBER: 138:385679

TITLE: Ammonium salts from polymer-bound N-hydroxysuccinimide

as solid-supported reagents for EDC-mediated

amidations

AUTHOR(S): Chinchilla, Rafael; Dodsworth, David J.; Najera,

Carmen; Soriano, Jose M.

CORPORATE SOURCE: Facultad de Ciencias, Departamento de Quimica

Organica, Universidad de Alicante, Alicante, 03080,

Spain

SOURCE: Tetrahedron Letters (2002), Volume Date 2003, 44(3), 463-466

CODEN: TELEAY; ISSN: 0040-4039

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 138:385679

New ammonium and alkylammonium salts derived from a polymeric N-hydroxysuccinimide (P-HOSu) have been prepared and used for the amidation

of carboxylic acids and amino acids mediated by 1-ethyl-3-[3-(dimethylamino)propyl]carbodiimide hydrochloride (EDC). These

polymer-supported ammonium salts afforded the corresponding amides in good yield, without detectable α-racemization and with easy recovery of

the P-HOSu after the amidation reaction, being especially suitable for the

amidation of Fmoc-protected amino acids.

REFERENCE COUNT: 54 THERE ARE 54 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L42 ANSWER 7 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2002:908277 CAPLUS

DOCUMENT NUMBER: 138:254580

TITLE: IBX-mediated oxidation of primary alcohols and

aldehydes to form carboxylic acids

AUTHOR(S): Mazitschek, Ralph; Mulbaier, Marcel; Giannis,

Athanassios CORPORATE SOURCE:

Institut fur Organische Chemie Universitat Leipzig,

Leipzig, 04103, Germany

SOURCE: Angewandte Chemie, International Edition (2002),

41(21), 4059-4061

CODEN: ACIEF5; ISSN: 1433-7851 Journal

PUBLISHER: Wiley-VCH Verlag GmbH

& Co. KGaA DOCUMENT TYPE:

LANGUAGE: English OTHER SOURCE(S): CASREACT 138:254580

AB Primary alcs. and aldehydes were oxidized by 1-hydroxy-1,2-benziodoxole-

3(1H) - one 1-oxide in presence of the O-nucleophiles 2-hydroxypyridine, 1-hydroxybenzotriazole, and N-hydroxysuccinimide (NHS) to give carboxylic acids. The NHS-mediate oxidation yielded active ester (0-succinimidyl) in

most cases. REFERENCE COUNT: 19 THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD, ALL CITATIONS AVAILABLE IN THE RE FORMAT

L42 ANSWER 8 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

2002:675821 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 137:222033

TITLE: Compositions and methods for enhancing drug delivery

across and into ocular tissues

English

INVENTOR(S): Rothbard, Jonathan B.; Wender, Paul A.; McGrane, P. Leo; Sista, Lalitha Vs; Kirschberg, Thorsten A.

PATENT ASSIGNEE(S):

Cellgate, Inc., USA PCT Int. Appl., 119 pp. SOURCE:

CODEN: PIXXD2 DOCUMENT TYPE: Patent

LANGUAGE: FAMILY ACC. NUM. COUNT: 4

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE A1 20020906 WO 2002-US5804 WO 2002067917 20020225 <--

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            CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
            GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
            LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
            PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ,
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                              20040102
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        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
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                                          JP 2002-567285
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PRIORITY APPLN. INFO.:
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                                           US 1999-150510P
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                                                              A2 20000824
                                           US 2000-648400
                                                              W 20020225
                                           WO 2002-US5804
                        MARPAT 137:222033
OTHER SOURCE(S):
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AB Compns. and methods for enhancing delivery of drugs, diagnostic and other

agents across epithelial tissues, including into and across ocular tissues and blood-brain barrier are provided. The compns. and methods employ a delivery enhancing transporter that has sufficient guanidino or amidino side chain moieties to enhance delivery of a compound conjugated to the reagent across one or more layers of the tissue, compared to the non-conjugated compound The delivery-enhancing polymers include, for example, poly-arginine mols. that are preferably between about 6 and 25 residues in length. For example, a series of structural characteristics including sequence length, amino acid composition, and chirality that influence the ability of Tat49-57 to enter cells is identified. These characteristics provided the blueprint for the design of a series of novel peptoids, of which 17 members were synthesized and assayed for cellular uptake. This research established that the peptide backbone and hydrogen bonding along that backbone are not required for cellular uptake, that the quanidino head group is superior to other cationic subunits, and most significantly, that an extension of the alkyl chain between the backbone and the head group provides superior transporters. In addition to better uptake performance, these novel peptoids offer several advantages over Tat49-57 including cost-effectiveness, ease of synthesis of analogs, and protease stability. These features along with their significant water solubility (>100 mg/mL) indicate that these novel peptoids could serve as effective transporters for the mol. delivery of drugs, drug candidates, and other agents into cells.

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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L42 ANSWER 9 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN
                        2002:505440 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                        137:58577
TITLE:
                        Photoactivatable nucleic acid derivatives, their
                        synthesis and use in preparing immobilized nucleic
                        acid arrays
INVENTOR(S):
                        Guire, Patrick E.; Swanson, Melvin J.; Opperman, Gary
                        W.
PATENT ASSIGNEE(S):
                        USA
SOURCE:
                        U.S. Pat. Appl. Publ., 11 pp., Cont.-in-part of U.S.
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Ser. No. 916,913. CODEN: USXXCO Patent

English

LANGUAGE: FAMILY ACC. NUM. COUNT: 3

DOCUMENT TYPE:

PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE

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| ΕP | 1577 | 670 | | | A2 | | 2005 | 0921 | EP | 2005-6595 | | | 19980 | 811 | |
| EP | 1577 | 670 | | | A3 | | 2005 | 1207 | | | | | | | |
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| WO | 9943 | 688 | | | A1 | | | | | | | | | | |
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- US 2000-591564 A1 20000609 AB A photoactivatable nucleic acid derivative composition in which one or more photoreactive group(s) are bound to a natural or synthetic nucleic acid is disclosed. The photoreactive groups may be a ketone such as benzophenone, or may be a group which generates a nitrene or carbene. The photoreactive groups can be bound to the nucleic acid before, during or after its formation, and can thereafter be activated in order to attach the nucleic acid to another mol., e.g., to the surface of a solid support. Also described is a method of preparing such a composition in which a nucleic acid derivative containing a thermochem. reactive group is reacted with a compound containing
 - a reactive group and a photoreactive group. For example, reactions between amines and N-oxysuccinimde esters, between carboxylic acid chlorides and amines, or between a maleimide and a sulfhydryl group may be used to prepare the photoactive nucleic acid derivative Alternatively, nucleotide monomers containing a photoreactive group may be used in synthesis of oligonucleotides/nucleic acids. Thus, N-[3-(4benzovlbenzamido)propvllmethacrvlamide (BBA-APMA) and N-succinimidvl 6-maleimidohexanoate (MAL-EAC-NOS) were synthesized and, using these compds., a copolymer of acrylamide, BBA-APMA, and MAL-EAC-NOS was also synthesized. An amino-terminated oligonucleotide was immobilized on polypropylene or polyvinyl chloride microwell plates by irradiation in the presence of this copolymer.

DOCUMENT NUMBER: 137:79227

TITLE: Novel functional peptide nucleic acid monomer and

process for producing the same

INVENTOR(S): Ikeda, Hisafumi; Saito, Isao; Kitagawa, Fumihiko

PATENT ASSIGNEE(S): Applied Biosystems Japan Ltd., Japan

GOURCE: PCT Int. Appl., 63 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: Japanese FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | KIND DATE | APPLICATION NO. | DATE |
|------------------------|------------------|-------------------------|-------------|
| | | | |
| WO 2002051797 | A1 20020704 | WO 2001-JP8120 | 20010919 < |
| W: JP, US | | | |
| RW: AT, BE, CH, | CY, DE, DK, ES, | FI, FR, GB, GR, IE, IT, | LU, MC, NL, |
| PT, SE, TR | | | |
| EP 1357112 | A1 20031029 | EP 2001-970133 | 20010919 < |
| R: AT, BE, CH, | DE, DK, ES, FR, | GB, GR, IT, LI, LU, NL, | SE, MC, PT, |
| IE, FI, CY, | TR | | |
| US 2004101839 | A1 20040527 | US 2003-250592 | 20031224 |
| US 7282575 | B2 20071016 | | |
| PRIORITY APPLN. INFO.: | | JP 2000-394669 | A 20001226 |
| | | WO 2001-JP8120 | W 20010919 |
| OTHER SOURCE(S): | CASREACT 137:792 | 27; MARPAT 137:79227 | |

GI

AR A peptide nucleic acid (PNA) monomer represented by the following general formula A-(CH2)nCO-B [I; wherein A = Q or Q1 (wherein X = OH, Z = O; X = NH2, Z = H2N+; or X = NMe2, Z = Me2N+), Q2, Q3, Q4 (wherein R = hydrogen, NO2, NH2, NHCbz, bromine, fluorine, chlorine, or SO3Na2), Q5, 3-(4-dimethylaminophenylazo)phenyl, 4-(4-dimethylaminophenylazo)phenylsulf onvlamino, 2-(4-hydroxyphenylazo)benzovlamino, 5dimethylaminonaphthalenesulfonylamino, 1-pyrenecarbonyl, 1-pyrenylmethyl, 1-pyrenesulfonylamino, 6,7,8-trimethyl-1,3-dioxo-2,5-dihydro-2,4diazaphenazin-2-yl, 4-methylcoumarin-7-ylaminocarbonyl, 4-trifluoromethylcoumarin-7-ylaminocarbonyl, 4-methyl-2-oxo-1,2dihydroquinoin-7-ylaminocarbonyl, 2-oxo-1,2-dihydroquinoin-3ylaminocarbonyl, etc.; B is OH, pentafluorophenyloxy, succinimidyloxy, N-carboxylmethyl-N-[2-(tert-butoxycarbonylamino)ethyl]amino; n = an integer of 1 to 4] is prepared A PNA monomer I [A, N = same as above; B = N-carboxylmethyl-N-[2-(tert-butoxycarbonylamino)ethyl]amino] is prepared by amidation of an active ester I (A, n = same as above; B = pentafluorophenyloxy, succinimidyloxy) with tertbutoxycarbonylaminoethylamine or an @-amino acid derivative, in particular 2-[N-[2-(tert-butoxycarbonylamino)ethyl]aminolacetic acid (II). This process is convenient for the preparation of a photofunctional PNA monomer which is unstable under alkali condition. Thus, to a solution of 100 mg 2-(5,7,8-trimethyl-1,3-dioxo-2,5-dihydro-2,4-diazaphenazin-2-yl)acetic acid and 70.2 mg pentafluorophenol in 10 mL DMF was added 73.2 mg 1-ethy1-3-(3-dimethylaminopropyl)carbodiimide hydrochloride (EDC) at 0° and stirred at 0° for 1 h and at room temperature for 12 h to give 85% 2,3,4,5,6-pentafluorophenyl 2-(5,7,8-trimethyl-1,3-dioxo-2,5dihydro-2,4-diazaphenazin-2-yl)acetate (III). To a solution of the active

^{*} STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

ester III (100 mg) and 45.4 mg II in 10 mL DMF was added 36.3 μ L diisopropylethylamine and stirred at room temperature for 15 h to give 85% 2-[N-[2-(tert-butoxycarbonylamino)ethyl]-2-[(5,7,8-trimethyl-1,3-dioxo-2,5-dihydro-2,4-diazaphenazin-2-yl)acetyl]amino]acetic acid.

REFERENCE COUNT: 14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L42 ANSWER 11 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2002:465982 CAPLUS

DOCUMENT NUMBER: 137:47213

TITLE: Preparation of fused pyrimidinones and

benzodioxaborolidinylpropylaminopyrrolo[1,2a]pyrimidines as inhibitors of hepatitis C ns3 protease for the treatment of hepatitis C and other

viral diseases

INVENTOR(S): Glunz, Peter W.; Douty, Brent D.; Han, Wei
PATENT ASSIGNEE(S): Bristol-Myers Squibb Pharma Company, USA

SOURCE: PCT Int. Appl., 270 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

| | PATENT NO. | | | | | | KIND DATE | | | APPLICATION NO. | | | | | | | | | |
|----------|--------------------------|--------------------------|--------------------------|--------------------------|--------------------------|-------------------------|---------------------------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|--|--|
| WO | 2002048116
2002048116 | | | | | | | | WO 2 | 001- | US47 | 911 | | 20011212 < | | | | | |
| | W: | CO,
GM,
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NE, | AT, | BE, | CH,
TR, | | |
| US | | | | | | | 20020624
20030403 | | | AU 2002-30763 | | 3
4
90P | 20013
20013 | | 0011 | 212 < | | | |
| OTHER SO | OTHER SOURCE(S): | | | | | PAT | 137: | 4721 | | WO 2 | 001- | US47 | 911 | | W 2 | 0011 | 212 | | |

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

carbocyclic ring, alkylidene] and particularly dioxaborolylpropylamino pyrrolopyriminecarboxamides such as II are prepared as inhibitors of hepatitis C viral protein ns3 protease for the treatment of hepatitis C and other viral diseases. E.g., esterification of L-pyroglutamic acid with AcOCM63 and HClo4, thionation with Lawesson's reagent, S-methylation with MeI, and amidation with NH4Cl gives nonracemic aminopyrrolinecarboxylate III. Treatment of III with di-Me 2-(methoxymethylene)malonate, hydrolysis of the Me ester moiety with LiOH, preparation of the acyl azide with diphenylphosphoryl azide and Curtius rearrangement in the presence of PhCH2OH, and hydrolysis of the tert-Bu ester with CF3CO2H gives pyrrolo[1,2-a]pyrimidine IV. Coupling of IV with an α -allyl aminomethylboronate pinanediol ester gives II. I inhibit hepatitis C ns3 protease with IC50 values of <100 μ M. Pharmaceutical compns. containing I are given.

L42 ANSWER 12 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2002:408568 CAPLUS

DOCUMENT NUMBER: 137:8158

TITLE: Manufacture and uses of Hollow Polymeric microspheres INVENTOR(S): Walt, David R.; Mandal, Tarun K.; Fleming, Michael S. PATENT ASSIGNEE(S): Tufts University. USA

SOURCE: PCT Int. Appl., 55 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

| PA | TENT : | NO. | KIND DATE | | | APPLICATION NO. | | | | | | DATE | | | | | |
|----------|---------------------------------------|---|---|---|---|---|--|------------------------------------|--------------------------|--------------------------|---|---|----------------------------|--------------------------|----------------------------------|--------------------------|--|
| | | | | | A2 20020530
A3 20030605 | | | WO 2001-US51278 | | | | | | 20011025 < | | | |
| wo | W: | AE,
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| US
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16 | LU,
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B2 | NL,
NE, | PT,
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2004 | SE,
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AB The invention features core-shell microsphere compns., hollow polymeric microspheres, and methods for making the microspheres. The microspheres are characterized as having a polymeric shell with consistent shell thickness. One method includes polymerizing one or more monomers or a polymerizable grafted unit over a solid inorg, core microsphere to form a shell, and then the solid core is etched away by acid. The solid inorg, core has had the polymer initiator previously attached to the surface. Another method mixes polymeric nanospheres over solid inorg, core microsphere, the mixture heated, and the solid core is etched away. The hollow microspheres can be filled with dye materials, therapeutic materials, or other substances.

L42 ANSWER 13 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2002:175766 CAPLUS

DOCUMENT NUMBER: 137:155063

TITLE: Evaluation of Morphogenic Regulatory Activity of Farnesoic acid and Its Derivatives against Candida

albicans Dimorphism

AUTHOR(S): Kim, Sanghee; Kim, Eunkvung; Shin, Dong-Sun; Kang,

Heonicong; Oh, Ki-Bong

CORPORATE SOURCE: Seoul National University, Natural Products Research

Institute, Seoul, Jongro, 110-460, S. Korea

SOURCE: Bioorganic & Medicinal

Chemistry Letters (2002), 12(6), 895-898

CODEN: BMCLE8; ISSN: 0960-894X

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 137:155063

A series of farnesoic acid derivs, was prepared and their morphogenic regulatory activities were evaluated. Their inhibitory activities against yeast cell growth and yeast-to-hypha transition examined in Candida albicans cells are dependent upon the chain length as well as the substitution patterns on the isoprenoid template. The preliminary structure-activity relationship of these compds. is described to elucidate the essential

structural requirements.

REFERENCE COUNT: THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L42 ANSWER 14 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2002:119134 CAPLUS

DOCUMENT NUMBER: 138:333559

TITLE: An Inhibitor of the Human UDP-GlcNAc 4-Epimerase

Identified from a Uridine-Based Library. A Strategy to

Inhibit O-Linked Glycosylation

AUTHOR(S): Winans, Katharine A.; Bertozzi, Carolyn R.

CORPORATE SOURCE: Department of Chemistry, Center for New Directions in

Organic Synthesis, University of California, Berkeley,

CA, 94720, USA Chemistry & Biology (

SOURCE:

2002), 9(1), 113-129

CODEN: CBOLE2: ISSN: 1074-5521

PUBLISHER: Elsevier Science Ltd.

54

DOCUMENT TYPE: Journal

LANGUAGE: English

CASREACT 138:333559 OTHER SOURCE(S):

The biol. study of O-linked glycosylation is particularly problematic, as chemical tools to control this modification are lacking. An inhibitor of the UDP-GlcNAc 4-epimerase that synthesizes UDP-GalNAc, the donor initiating O-linked glycosylation, would be a powerful reagent for reversibly inhibiting O-linked glycosylation. We synthesized a 1338 member library of uridine analogs directed to the epimerase by virtue of substrate mimicry. Screening of the library identified an inhibitor with a Ki value of 11 µM. Tests against related enzymes confirmed the compound's specificity for the UDP-GlcNAc 4-epimerase. Inhibitors of a key step of O-linked glycan biosynthesis can be discovered from a directed library screen. Progeny thereof may be powerful tools for controlling O-linked glycosylation in cells.

REFERENCE COUNT:

THERE ARE 54 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT L42 ANSWER 15 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2001:905597 CAPLUS

DOCUMENT NUMBER: 136:263357

TITLE: Nucleosides derived from urocanic acid: potential

ligands for CG base pairs

AUTHOR(S): Purwanto, Maria G. M.; Lengeler, David; Weisz, Klaus CORPORATE SOURCE: Institut fur Chemie der Freien Universitat Berlin,

Berlin, D-14195, Germany

SOURCE: Tetrahedron Letters (2001), Volume Date 2002, 43(1),

61-64

CODEN: TELEAY; ISSN: 0040-4039

PUBLISHER: Elsevier Science Ltd. DOCUMENT TYPE: Journal

LANGUAGE . English

OTHER SOURCE(S): CASREACT 136:263357

AB A nucleoside analog based on imidazole-4-acrylamide (urocanamide) was synthesized and studied for its use as a specific ligand for a

cytosine-guanosine Watson-Crick base pair. One- and two-dimensional 1H NMR expts. in methylene chloride at ambient and low temps. not only

indicate the strength of association but also confirm specific binding of the novel nucleoside to the base pair through the formation of two hydrogen

bonds.

REFERENCE COUNT: 19 THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L42 ANSWER 16 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2001:863431 CAPLUS DOCUMENT NUMBER: 136:2448

TITLE: Sensor for analyte detection

INVENTOR(S): Bauer, Alan Joseph

PATENT ASSIGNEE(S): Biosensor Systems Design., Inc., USA SOURCE: U.S., 25 pp., Cont.-in-part of U.S. 6,096,497.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

| PATENT NO. | | | | | KIND DATE | | | APPLICATION NO. | | | | | DATE | | | | | |
|-------------|------------|---------------------------------|--------------------------|---------------------------------|---------------------------------|--|---------------------------------|--------------------------------------|---|--------------------------|----------------------------------|--------------------------|---------------------------------|--------------------------|--------------------------|--------------------------|--------------------------|---|
| US | US 6096497 | | | | | A 20000801 | | | US 2000-701906
US 1998-110686
WO 1999-IL309 | | | | | | 19980707 < | | | < |
| | W: | AE,
DE,
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| DD 7 OD 7 W | | GH,
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SN, | NL,
TD, | PT,
TG | SE, | BF, | BJ, | CF, | CG, | |
| PRIORITY | | | | | | IL 1:
US 1:
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IL 1:
WO 1:
IS 1: | 998-
998-
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999- | 1106
1257
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1L30 | 86
20
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54 | | A2 1
A 1
A 1
A 1
W 1 | | 707
811
112
504
610 | | | | | |

AB A sensor for detecting analytes is described. Analyte presence or concentration

is determined through measurement of changes in induced electromotive force. current or other

elec. property in a base member during analyte exposure to the sensor. According to one class of embodiments, the present device immobilizes natural or synthetic macromols, sufficiently close to an elec.-conductive base member to insure that any alteration in the motion and/or

electrostatic fields of the macromols, during interaction with a predetd. analyte will induce an increased electromotive force in the base member.

REFERENCE COUNT: 24 THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L42 ANSWER 17 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2001:598434 CAPLUS

DOCUMENT NUMBER: 135:177719

TITLE: Target molecule attachment to surfaces

INVENTOR(S): Chappa, Ralph A.; Hu, Sheau-Ping; Swan, Dale G.;

Swanson, Melvin J.; Guire, Patrick E.

PATENT ASSIGNEE(S): Surmodics, Inc., USA SOURCE: U.S. Pat. Appl. Publ., 26 pp., Cont.-in-part of U.S.

5,858,653. CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 5

PATENT INFORMATION:

| PA | TENT I | NO. | | | KIN |) | DATE | | | APP | LIC | CAT: | ION I | NO. | | | DA | TE | | |
|---------------|--------------------------|-----|------|-----|-----|----------|----------|------|-----|----------------|------|------|-------|-----|-----|----|------------|------|-----|---|
| | 2001 | | | | | | 2001 | | | US | 199 | 99-2 | 2279 | 13 | | | 19 | 990 | 108 | < |
| | US 6465178
US 5858653 | | | | | | 19990112 | | | US 1997-940213 | | | | | | | 19970930 < | | | |
| | CA 2360000 | | | | | | 20000713 | | | | | | | | | | | | | |
| | WO 2000040593 | | | | | | 20000713 | | | | | | | | | | | | | |
| | WO 2000040593 | | | | | | | | | | | | | | | | | | | |
| | W: | AU, | CA, | JP, | MX | | | | | | | | | | | | | | | |
| | RW: | AT, | BE, | CH, | CY, | DE, | DK, | ES, | FI, | FF | ٠, ٥ | GΒ, | GR, | IE, | IT, | LU | , | MC, | NL | , |
| | | PT, | SE | | | | | | | | | | | | | | | | | |
| EP | 1141 | | | | | | | | | | | | | | | | | | | |
| | R: | ΑT, | BE, | CH, | DE, | DK, | ES, | FR, | GB, | GF | ٠, : | ΙT, | LI, | LU, | NL, | SE | , | MC, | PT | , |
| | | IE, | | | | | | | | | | | | | | | | | | |
| | 2002 | | | | | | 2002 | | | | | | | | | | | | | < |
| AU | AU 778265 | | | | B2 | 20041125 | | | | | | | | | | | | | | |
| US 2003113792 | | | | | | 2003 | | | US | 200 | 00- | 5215 | 45 | | | 20 | 000 | 309 | < | |
| | 6762 | | | | B2 | | 2004 | | | | | | | | | | | | | |
| | 2001 | | | | | | 2001 | | | | | | | | | | | | | < |
| | 2003 | | | | | | 2003 | | | | | | | | | | | | | < |
| | 2004 | | | | | | 2004 | | | US | 200 | 04-1 | 3446 | 67 | | | 20 | 040 | 512 | |
| | 7300 | | | | | | 2007 | | | | | | | | | | | | | |
| | 2005 | | | | A1 | | 2005 | 0804 | | | | | | 71 | | | | | | |
| PRIORIT | Y APP | LN. | INFO | . : | | | | | | | | | | 13 | | | | 970 | | |
| | | | | | | | | | | | | | | 13 | | | | 990 | | |
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| | | | | | | | | | | US | 200 | 02-3 | 1929 | 17 | | A3 | 20 | 020 | 709 | |

AB Method and reagent composition for covalent attachment of target mols., such as nucleic acids, onto the surface of a substrate are described. The reagent composition includes groups capable of covalently binding to the target mol. Optionally, the composition can contain photoreactive groups for use in

he

attaching the reagent composition to the surface. The reagent composition can

used to provide activated slides for use in preparing microarrays of nucleic acids. Glass slides coated with a copolymer of acrylamide,

N-[3-(4-benzoylbenzamido)propyl]methacrylamide (BBA-APMA), and

N-succinimidyl 6-maleimidohexanoate (MAL-EAC-NOS) (preparation given) were reacted with amine-modified PCR products from the β -galactosidase

gene using microarraving spotting pins.

L42 ANSWER 18 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2001:565638 CAPLUS

DOCUMENT NUMBER: 135:266385

TITLE: 4:3-β-Naphthapyrone-4-acetic acid

N-hydroxy-succinimidyl ester as a fluorescent labeling

reagent for amino acids and oligopeptides in high-performance liquid chromatography

AUTHOR(S): Liu, Xin; Wang, Hong; Liang, Shu-Cai; Zhang, Hua-Shan

Department of Chemistry, Wuhan University, Wuhan, CORPORATE SOURCE:

430072, Peop. Rep. China

SOURCE: Chromatographia (2001), 53(5/6), 326-330

CODEN: CHRGB7; ISSN: 0009-5893

Friedrich Vieweg

& Sohn Verlagsgesellschaft mbH DOCUMENT TYPE: Journal LANGUAGE: English

4:3-β-Naphthapyrone-4-acetic acid N-hydroxysuccinimidyl ester

(NPA-OSu) is a highly sensitive and moderately reactive derivatizing

reagent with a naphthapyrone moiety as fluorophore and an N-hydroxysuccinimidyl active ester as reactive group toward amino compds.

It is readily prepared in two steps. The fluorescence properties of NPA-OSu and its hydrolysis product were studied in detail, and the conditions for derivatization and separation of the NPA-OSu derivs. of some amino acids and oligopeptides were studied. At $\lambda ex = 352$ nm and $\lambda em = 422$

nm the detection limits (signal-to-noise ratio = 3) for amino acids and

oligopeptides reached fmol levels for injection of 20 µL; this sensitivity was comparable with that obtained using 7-

(diethylamino)coumarin-3-carboxylic acid succinimidyl ester as

derivatizing reagent in the anal. of amino acids by capillary electrophoresis with laser-induced-fluorescence detection.

REFERENCE COUNT: 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L42 ANSWER 19 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2001:241696 CAPLUS

DOCUMENT NUMBER: 134:265244

TITLE: Antibody catalysis of enantio- and diastereo-selective aldol reactions

Barbas, Carlos F.; Lerner, Richard A.; Zhong, Guofu INVENTOR(S):

PATENT ASSIGNEE(S): The Scripps Research Institute, USA U.S., 15 pp., Cont. of U.S. Ser. No. 415,453.

SOURCE: CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------|------|----------|-----------------|------------|
| | | | | |
| US 6210938 | B1 | 20010403 | US 1999-458367 | 19991209 < |
| US 6294374 | B1 | 20010925 | US 1999-415453 | 19991008 < |

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CA 2389250 A1 20010419 CA 2000-2389250 20001006 <--
WO 2001027145 A1 20010419 WO 2000-US27777 20001006 <--
          W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
               CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
                HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
                LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
                SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU,
           RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
                DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ,
                CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                              A1 20020807 EP 2000-968865
          R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
                IE, SI, LT, LV, FI, RO, MK, CY, AL
      US 2001018201 A1 20010830
US 6309881 B2 20011030
                                                  US 2001-824279
                                                                                20010402 <--
      US 6309881
                                                    US 1999-415453 A1 19991008
US 1999-458367 A 19991209
WO 2000-US27777 W 20001006
PRIORITY APPLN. INFO.:
OTHER SOURCE(S):
                             CASREACT 134:265244; MARPAT 134:265244
AB Nine efficient aldolase antibodies were generated using a sulfone
      β-diketone hapten. This hapten combines, in a single mol.,
      structural components employed for reactive immunization with structural
      components employed for forming a transition state analog of the aldol
      reaction. Characterization of 2 of these antibodies reveals that they are
      highly proficient (≤1000-fold better than any other antibody
      catalyst) and enantioselective catalysts for aldol and retro-aldol
      reactions and exhibit enantio- and diastereo- selectivities opposite that
      of antibody 38C2.
REFERENCE COUNT: 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS
                                    RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L42 ANSWER 20 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER: 2001:221918 CAPLUS
DOCUMENT NUMBER:
                             134:249193
TITLE:
                             Test kit and electrode sensor for multi-array,
                             multi-specific electrochemiluminescence testing
INVENTOR(S):
                             Wohlstadter, Jacob N.; Wilbur, James; Sigal, George;
                             Martin, Mark; Guo, Liang-Hong; Fischer, Alan; Leland,
                             Jon: Billadeau, Mark A.
PATENT ASSIGNEE(S):
                            Meso Scale Technologies, LLC, USA
SOURCE:
                             U.S., 103 pp., Cont.-in-part of U.S. 6,066,448.
                             CODEN: USXXAM
DOCUMENT TYPE:
                             Patent
LANGUAGE:
                             English
FAMILY ACC. NUM. COUNT: 6
PATENT INFORMATION:
     PATENT NO. KIND DATE APPLICATION NO. DATE

US 6207369 B1 20010327 US 1996-715163 19960917 <--
US 6066448 A 20000523 US 1996-611804 19960306 <--
CN 1661115 A 20050831 CN 2005-10005720 19960306
ZA 9601925 A 19970805 ZA 1996-1925 19960308 <--
US 6140045 A 20001031 US 1997-814085 19970306 <--
CA 2265828 A1 19980326 W 1997-US 1997-285828 19970917 <--
W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE,
W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE,
W: BE EF EF EF GR GR GH HII TO IL LS JE KE KG KP KR
               DK, EE, ES, FI, GB, GE, GH, HU, ID, IL, IS, JP, KE, KG, KP, KR,
                KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ,
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PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG,
              UZ, VN, YU, ZW
          RW: GH, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI, FR,
              GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA,
              GN, ML, MR, NE, SN, TD, TG
     AU 9746495
                           A
                                   19980414
                                               AU 1997-46495
                                                                          19970917 <--
     AU 743567
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                                   20020131
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ZA 9708380
                                  19980417 ZA 1997-8380
19990929 EP 1997-945249
                           A
                                                                          19970917 <--
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                            A1
                                                                         19970917 <--
          R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
              IE, FI
     JP 2001503856
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                                  20010321
                                                JP 1998-514984
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                                                                          19971017 <--
                         A 20000626 KR 1999-702230
A1 20010913 US 2001-771796
A 20020523 AU 2002-29296
A1 20040506 US 2003-693441
B2 20060321
     KR 2000036176
                                  20000626 KR 1999-702230
                                                                          19990316 <--
     US 2001021534
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     AU 200229296
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US 7015046 B2 20060321
AU 2005201886 A1 20050526
AU 2005201886 B2 20070906
JP 2006047321 A 20060216
US 2006068499 A1 20060330
US 2006172340 A1 2006080
PRIORITY APPLN, INFO.;
                                               AU 2005-201886
                                                                          20050504
                                               JP 2005-296368
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                                                US 2005-264535
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                                                US 2005-300808
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                                                                    B2 19950310
B2 19950310
A2 19960306
                                                 US 1995-402076
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                                                 US 1996-611804
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                                                                     A3 19960306
                                                 JP 1996-527737
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                                                 US 1996-12957P
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                                                 WO 1996-US3190
                                                                     A 19960306
                                                 US 1996-715163 A 19960917
                                                                     A3 19970917
                                                 US 1997-932110
                                                 WO 1997-US16942
                                                                     W 19970917
                                                 US 2001-771796
                                                                     B1 20010129
                                                 AU 2002-29296
                                                                      A3 20020328
                                                 US 2003-693441 A1 20031024
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AB Materials and methods are provided for producing patterned multi-array, multi-sp. surfaces for use in diagnostics. The invention provides for electrochemiluminescence methods for detecting or measuring an analyte of interest. It also provides for novel electrodes for ECL assays. Materials and methods are provided for the chemical and/or phys. control of conducting domains and reagent deposition for use multiply specific testing procedures. An ECL immunoassay for TSR used a composite electrode of EVA and carbon fibrils. A DNA hybridization assay was performed on a fibril-polymer composite.

REFERENCE COUNT: 82 THERE ARE 82 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L42 ANSWER 21 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2000:838129 CAPLUS

DOCUMENT NUMBER: 134:5118

TITLE: Derivatized oligonucleotides having improved uptake

and other properties
INVENTOR(S): Manoharan, Muthiah; Cook, Phillip Dan; Bennett,

Clarence Frank

PATENT ASSIGNEE(S): ISIS Pharmaceuticals, Inc., USA

SOURCE: U.S., 25 pp., Cont.-in-part of U.S. Ser. No. 782,374, abandoned.

CODEN: USXXAM

DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 326

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE WU 6153737 A 20001128 US 1994-211882 WO 9110671 A1 19910725 WO 1991-US243 19940422 <--19910111 <--W: AU, BR, CA, FI, HU, JP, KR, NO, US RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LU, NL, SE EP 1418179 A2 20040512 EP 2003-78862 EP 1418179 A3 20060308 19910111 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE CA 2089376 A1 19920214 CA 1991-2089376 19910812 <--EP 1443051 A2 20040804 EP 2004-76246 19910812 EP 1443051 A3 20050817 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE AT 318273 T 20060315 AT 1991-915355 19910812 ES 2259177 T3 20060916 ES 1991-915355 19910812 W0 9307883 A1 19930429 W0 1992-U99196 19921023 19921023 <--W: AU, BB, BG, BR, CA, CS, FI, HU, JP, KP, KR, LK, MG, MN, MW, NO, PL. RO. RU. US RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, SN, TD, TG EP 1331011 A2 20030730 EP 2003-76286 EP 1331011 A3 20031217 20031217 EP 1331011 A3 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, IE US 5578718 A 19961126 US 1993-116801 19930903 <--US 1990-463358 B2 19900111
US 1990-666977 B2 19900813
WO 1991-105243 A2 19910111
US 1991-782374 B2 19911024
WO 1992-US9196 W 19921023
BP 1991-903066 A3 19910111
BP 1991-915355 A3 19910812
EP 1992-922139 A3 19921023
AU 1993-38025 A3 1992023
US 1993-116801 A2 1993093
US 1994-211882 A2 19940422
US 1995-458396 A1 1995002
US 1995-948151 A1 19971009
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US 2002-73718 A1 20020211 US 2002-154993 A1 20020523

AB Linked nucleosides having at least one functionalized nucleoside that bears a substituent such as a steroid mol., a reporter mol., a non-aromatic lipophilic mol., a reporter enzyme, a peptide, a protein, a water soluble vitamin, a lipid soluble vitamin, an RNA cleaving complex, a metal chelator, a porphyrin, an alkylator, a pyrene, a hybrid photo-nuclease/intercalator, or an arvl azide photo-crosslinking agent exhibit increased cellular uptake and other properties. The substituent can be attached at the 2'-position of the functionalized nucleoside via a linking group. If at least a portion of the remaining linked nucleosides are 2'-deoxy-2'-fluoro, 2'-0-methoxy, 2'-0-ethoxy, 2'-0-propoxy, 2'-O-aminoalkoxy or 2'-O-allyloxy nucleosides, the substituent can be attached via a linking group at any of the 3' or the 5' positions of the nucleoside or on the heterocyclic base of the nucleoside or on the inter-nucleotide linkage linking the nucleoside to an adjacent nucleoside.

REFERENCE COUNT: 36 THERE ARE 36 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L42 ANSWER 22 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2000:433344 CAPLUS

DOCUMENT NUMBER: 133:79341

TITLE: Immunostimulating and vaccine compositions employing saponin analog adjuvants and uses thereof

INVENTOR(S): Marciani, Dante J.

PATENT ASSIGNEE(S): Galenica Pharmaceuticals, Inc., USA

SOURCE: U.S., 40 pp., Cont.-in-part of U.S. 5,977,081.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|------|----------|------------------|------------|
| | | | | |
| US 6080725 | A | 20000627 | US 1999-290606 | 19990413 < |
| US 5977081 | A | 19991102 | US 1998-81647 | 19980520 < |
| PRIORITY APPLN. INFO.: | | | US 1997-47129P P | 19970520 |
| | | | US 1998-80389P P | 19980402 |
| | | | US 1998-81647 A2 | 19980520 |

OTHER SOURCE(S): MARPAT 133:79341

AB The present invention is directed to vaccines comprising (1) one or more bacterial, viral or tumor-associated antigens; and (2) one or more saponin-lipophile conjugate in which a lipophilic moiety such as a lipid, fatty acid, polyethylene glycol or terpene is covalently attached to a non-acvlated or desacvlated triterpene saponin via a carboxyl group present on the 3-O-glucuronic acid of the triterpene saponin. The attachment of a lipophile moiety to the 3-O-glucuronic acid of a saponin such as Quillaja desacylsaponin, lucyoside P, or saponin from Gypsophila, Saponaria and Acanthophyllum enhances their adjuvant effects on humoral and cell-mediated immunity. Addnl., the attachment of a lipophile moiety to the 3-0-qlucuronic acid residue of non- or des-acylsaponin yields a saponin analog that is easier to purify, less toxic, chemical more stable, and possesses equal or better adjuvant properties than the original saponin.

REFERENCE COUNT: 46 THERE ARE 46 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L42 ANSWER 23 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2000:117021 CAPLUS

DOCUMENT NUMBER: 132:166361

TITLE: Saturated and unsaturated abietane derivatives,

derived conjugates and uses in a diagnostic

composition, a reagent and a device

INVENTOR(S): Charles, Marie-helene; Piga, Nadia; Battail-Poirot,

Nicole; Veron, Laurent; Delair, Thierry; Mandrand,

PATENT ASSIGNEE(S): Bio Mer.

PATENT ASSIGNEE(S): Bio Merieux, Fr. SOURCE: PCT Int. Appl., 54 pp.

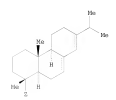
CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: French

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PA | TENT | NO. | | | KIN | | DATE | | | | ICAT | | | | | ATE | | |
|---------|-------|------|------|-----|-----|-----|------|------|-----|------|------|------|-----|-----|-----|------|-------|---|
| WO | 2000 | 0079 | 82 | | | | 2000 | 0217 | | | | | | | | 9990 | 727 | < |
| | W: | ΑE, | AL, | AM, | ΑT, | AU, | AZ, | BA, | BB, | BG, | BR, | BY, | CA, | CH, | CN, | CU, | CZ, | |
| | | DE, | DK, | EE, | ES, | FI, | GB, | GD, | GE, | GH, | GM, | HR, | HU, | ID, | IL, | IN, | IS, | |
| | | JP, | KE, | KG, | KP, | KR, | KZ, | LC, | LK, | LR, | LS, | LT, | LU, | LV, | MD, | MG, | MK, | |
| | | MN, | MW, | MX, | NO, | NZ, | PL, | PT, | RO, | RU, | SD, | SE, | SG, | SI, | SK, | SL, | ΤJ, | |
| | | TM, | TR, | TT, | UA, | UG, | US, | UZ, | VN, | YU, | ZA, | ZW, | AM, | ΑZ, | BY, | KG, | ΚZ, | |
| | | MD, | RU, | ΤJ, | TM | | | | | | | | | | | | | |
| | RW: | GH, | GM, | KE, | LS, | MW, | SD, | SL, | SZ, | UG, | ZW, | ΑT, | BE, | CH, | CY, | DE, | DK, | |
| | | | | | | | IE, | | | | | | SE, | BF, | ВJ, | CF, | CG, | |
| | | CI, | CM, | GA, | GN, | GW, | ML, | MR, | NE, | SN, | TD, | TG | | | | | | |
| FR | 2781 | 802 | | | A1 | | 2000 | 0204 | | FR 1 | 998- | 1008 | 4 | | 1 | 9980 | 731 - | < |
| FR | 2781 | 802 | | | B1 | | 2001 | 0511 | | | | | | | | | | |
| CA | 2339 | 102 | | | A1 | | 2000 | 0217 | | CA 1 | 999- | 2339 | 102 | | 1 | 9990 | 727 | < |
| | 9949 | | | | | | 2000 | 0228 | | AU 1 | 999- | 4917 | 3 | | 1 | 9990 | 727 | < |
| EP | 1100 | 773 | | | A1 | | 2001 | 0523 | | EP 1 | 999- | 9329 | 81 | | 1 | 9990 | 727 | < |
| | R: | AT, | BE, | CH, | DE, | DK, | ES, | FR, | GB, | GR, | IT, | LI, | LU, | NL, | SE, | MC, | PT, | |
| | | IE, | FΙ | | | | | | | | | | | | | | | |
| US | 2002 | 1554 | 96 | | A1 | | 2002 | 1024 | | US 2 | 001- | 7715 | 54 | | 2 | 0010 | 130 | < |
| PRIORIT | Y APP | LN. | INFO | . : | | | | | | | 998- | | | | | | | |
| | | | | | | | | | | WO 1 | 999- | FR18 | 46 | | W 1 | 9990 | 727 | |
| OTHER S | OURCE | (S): | | | MAR | PAT | 132: | 1663 | 61 | | | | | | | | | |



AB $\,$ The invention concerns a saturated or unsatd. abietane derivative (I) [Z = -COOR5,

-CONR1R2, -COONR3R4, -COR6, -CON, -COOR5, -CHOHR7, -SR8, -OR8, -CN, -CNO, -CNS, -NCO, -NCS, -R1R2CR9; R1, R2, R3, R4 = H, alkyl, (un)substituted

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aryl; alkene; alkyne; (un)substituted aminoacyl, (un)substituted peptidyl;
    R1, R2, or R3, R4 together can form a cycle or a heterocycle; R5 = H,
     alkyl, alkene, alkyne; aryl (un)substituted into C6-C20; R6 = H, halogen,
    alkyl, alkene, alkyne, aryl (un) substituted into C6-C20; R7, R8 = H,
    alkyl, alkene, alkyne; R9 = -CN, -CNO, -CNS, -NCO, -NCS]. The invention
    also concerns a derived conjugate with oligonucleotide, anti-alpha
     fetoprotein, oligomer or bovine serum albumin and the use of said derivative
    and said conjugate in a diagnostic composition, a reagent and a device.
REFERENCE COUNT:
                        5
                             THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS
                              RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L42 ANSWER 24 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                       1999:811446 CAPLUS
DOCUMENT NUMBER:
                        132:47205
TITLE .
                        A sensor for analyte detection
INVENTOR(S):
                       Bauer, Alan Joseph
                       Biosensor Systems Design, Inc. (1998), USA
PATENT ASSIGNEE(S):
SOURCE:
                        PCT Int. Appl., 66 pp.
                        CODEN: PIXXD2
DOCUMENT TYPE:
                        Patent
LANGUAGE:
                        English
FAMILY ACC. NUM. COUNT: 3
PATENT INFORMATION:
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| | PATENT NO. | | | | | | | | APPLICATION NO. | | | | | | | | |
|---------|------------|------|------|--------|-------|------|------|------|-----------------|------|-----|-----|-----|-----|------|------|-------|
| | | | | | A1 | | | | | | | | | | | 9990 | 610 < |
| | W: | ΑE, | AL, | AM, | AT, | AU, | AZ, | BA, | BB, | BG, | BR, | BY, | CA, | CH, | CN, | CU, | CZ, |
| | | DE, | DK, | EE, | ES, | FI, | GB, | GD, | GE, | GH, | GM, | HR, | HU, | ID, | IL, | IN, | IS, |
| | | JP, | KΕ, | KG, | KP, | KR, | ΚZ, | LC, | LK, | LR, | LS, | LT, | LU, | LV, | MD, | MG, | MK, |
| | | MN, | MW, | MX, | NO, | ΝZ, | PL, | PΤ, | RO, | RU, | SD, | SE, | SG, | SI, | SK, | SL, | TJ, |
| | | | | | | UG, | US, | UZ, | VN, | YU, | ZA, | ZW, | ΑM, | ΑZ, | BY, | KG, | KZ, |
| | | | RU, | | | | | | | | | | | | | | |
| | RW: | | | | | | SD, | | | | | | | | | | |
| | | | | | | | IE, | | | | | | SE, | BF, | ВJ, | CF, | CG, |
| | | | | | | | ML, | | | | | | | | | | |
| | 6096 | | | | | | | | | | | | | | | | 707 < |
| | | | | | | | | | | | | | | | | | 610 < |
| EP | | | | | | | | | | | | | 62 | | 1 | 9990 | 610 < |
| | | | | | | | FR, | | | | | | | | | | |
| | | | | | | | 2001 | 1127 | | | | | | | | | 205 < |
| PRIORIT | Y APP | LN. | INFO | . : | | | | | | | | | | | A 1 | | |
| | | | | | | | | | | US 1 | | | | | A2 1 | | |
| | | | | | | | | | | IL 1 | | | | | A 1 | | |
| | | | | | | | | | | IL 1 | | | | | A 1 | | |
| | | | | | | | | | | IL 1 | | | | | A 1 | | |
| | | | | | | | | | | | | | | | A 1 | | |
| | | | | | | | | | | | | | | | W 1 | | 610 |
| AR A | senso | r fo | r de | t.ect. | ina : | anal | vtes | of | | | | | | | | | 010 |

A sensor for detecting analytes of interest is described. Analyte presence or concentration is determined through measurement of changes in induced

electromotive force, current or other elec. property in a base member during analyte

exposure to the sensor. According to one class of embodiments, the present device immobilizes natural or synthetic macromols. sufficiently close to an elec.-conductive base member to insure that any alteration in the motion and/or electrostatic fields of the macromols. during interaction with a predetd. analyte induces an altered electromotive force in

the base member. In one example, the sensor described is used for food inspection. REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L42 ANSWER 25 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1999:45046 CAPLUS

DOCUMENT NUMBER: 130:121859

TITLE: Reagent having attracting and reacting groups for attaching target molecules to a surface

INVENTOR(S): Duran, Lise W.; Swanson, Melvin J.; Amos, Richard A.;

Hu, Sheau-ping J.; Guire, Patrick E.

PATENT ASSIGNEE(S): Surmodics, Inc., USA SOURCE: U.S., 19 pp.

CODEN: USXXAM
DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 5

PATENT INFORMATION:

| | TENT 1 | | | | | | DATE | | | | | | | DATE | |
|----------|----------------|------|------|-----|-----|-----|------|------|-------|--------|--------|-------|--------|--------|------|
| US | 58586 | 553 | | | A | | 1999 | 0112 | US | 1997- | 94021 | 3 | | 199709 | 30 < |
| CA | 23043 | 362 | | | A1 | | 1999 | 0408 | CF | 1998- | -23043 | 62 | | 199809 | 25 < |
| WO | 99169 | 907 | | | A2 | | 1999 | 0408 | WC | 1998- | US201 | 40 | | 199809 | 25 < |
| WO | 99169 | 907 | | | A3 | | 1999 | 0819 | | | | | | | |
| | W: | AU, | CA, | JP, | MX | | | | | | | | | | |
| | RW: | AT, | BE, | CH, | CY, | DE, | DK, | ES, | FI, E | R, GB, | GR, | IE, I | IT, LU | J, MC, | NL, |
| | | | SE | | | | | | | | | | | | |
| AU | 98958
73739 | 328 | | | A | | 1999 | 0423 | ΑU | 1998- | 95828 | | | 199809 | 25 < |
| AU | 73739 | 91 | | | B2 | | 2001 | 0816 | | | | | | | |
| | 10194 | | | | | | 2000 | | | 1998- | 94952 | 4 | | 199809 | 25 < |
| EP | 10194 | 124 | | | B1 | | 2005 | 0112 | | | | | | | |
| | | | | | GB, | | | | | | | | | | |
| JP | 20015 | 5186 | 04 | | T | | | | | | | | | 199809 | |
| | 20010 | | | | | | 2001 | 0816 | US | 1999- | -22791 | 3 | | 199901 | 08 < |
| US | 64651 | 178 | | | B2 | | 2002 | | | | | | | | |
| US | 20031 | 1137 | 92 | | A1 | | 2003 | | | 2000- | -52154 | 5 | | 200003 | 09 < |
| US | 67620 | 019 | | | B2 | | 2004 | | | | | | | | |
| | 20000 | | | | | | 2000 | | | | | | | 200003 | |
| | 20031 | | | | | | 2003 | | | | | | | 200207 | |
| | 20042 | | | | | | 2004 | | | 2004- | -84466 | 7 | | 200405 | 12 |
| | 7300 | | | | | | 2007 | | | | | | | | |
| | 20051 | | | | A1 | | 2005 | 0804 | | | | | | 200504 | |
| PRIORIT: | Y APPI | N. | INFO | . : | | | | | | | | | | 199709 | |
| | | | | | | | | | | | | | | 199809 | |
| | | | | | | | | | | | | | | 199901 | |
| | | | | | | | | | | 2000- | | | | 200003 | |
| | | | | | | | | | | 2002- | | | | 200207 | |

AB Disclosed are a method and reagent composition for covalent attachment of target mols., such as nucleic acids, onto the surface of a substrate. The reagent composition includes groups capable of attracting the target mol. as well as groups capable of covalently binding to the target mol., once attracted. Optionally, the composition can contain photoreactive groups for use in attaching the reagent composition to the surface. Microwell plates coated and photoreacted with a prepared random copolymer of acrylamide, N-[3-(4-benzoylbenzamido)propyl]methacrylamide (preparation given), N-succinimidyl 6-methacrylamidobeanoate (preparation given), and

^{[3-(}methacryloylamino)propyl]trimethylammonium chloride, provided significant binding of a 50-mer capture probe and good hybridization signals.

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L42 ANSWER 26 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1998:774292 CAPLUS

DOCUMENT NUMBER: 130:22307

TITLE: (Aminostyryl)pyridinium compounds for radiolabeling

cell membranes, and preparation thereof

INVENTOR(S): Lambert, Carol; Mease, Ronnie C.; McAfee, John G.

PATENT ASSIGNEE(S): Research Corporation Technologies, Inc., USA

SOURCE: U.S., 13 pp.
CODEN: USXXAM

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|--------|-----------|-----------------|------------|
| | | | | |
| US 5840859 | A | 19981124 | US 1996-673798 | 19960627 < |
| PRIORITY APPLN. INFO.: | | | US 1996-673798 | 19960627 |
| OTHER SOURCE(S): | MARPAT | 130:22307 | | |

GI

AB Compds. I (n = 4-16; Det = organic group comprising radioisotope or capable of chelating radioisotope; Z- = 1 equivalent of biol. acceptable anion) are provided. I are useful to radiolabel cellular membranes, as of hematopoietic cells. I are preferably employed in vitro, in combination with a pharmaceutically acceptable carrier or vehicle, to label populations of mammalian cells, such as blood cells, including mixed leukocytes or lymphocytes. When introduced into a mammalian host, such as a human patient or animal, the labeled cells such as the leukocytes or lymphocytes, localize at a site of inflammation, infection, malignancy, or the like, thus enabling the imaging of the site, for diagnostic purposes or to enable the effective targeting of therapeutic agents.

REFERENCE COUNT: 25 THERE ARE 25 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L42 ANSWER 27 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1998:197670 CAPLUS

DOCUMENT NUMBER: 128:254896

TITLE: Multi-array, multi-specific electrochemiluminescent testing

INVENTOR(S): Wohlstadter, Jacob N.; Wilbur, James; Sigal, George;

Martin, Mark; Guo, Liang-Hong; Fischer, Alan; Leland, Jon; Billadeau, Mark A.; Helms, Larry R.; Darvari,

Ramin
PATENT ASSIGNEE(S): Meso Scale

PATENT ASSIGNEE(S): Meso Scale Technologies, LLC, USA

SOURCE: PCT Int. Appl., 288 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: 6

PATENT INFORMATION:

| | | TENT : | | | | | | | | | | LICAT | | | | | ATE | | |
|-------|------|--------|------|------|-----|-----|------|------|------|-----|------|----------------|------|-----|-----|--------------|------|-----|---|
| | | 9812 | 539 | | | A1 | | 1998 | 0326 | | WO : | 1997- | US16 | 942 | | 1 | | | < |
| | | W: | AL, | AM, | ΑT, | ΑU, | ΑZ, | BA, | BB, | ВG, | BR, | BY, | CA, | CH, | CN, | CU, | CZ, | DE, | |
| | | | DK, | EE, | ES, | FI, | GB, | GE, | GH, | HU, | ID, | IL, | IS, | JP, | KE, | KG, | KP, | KR, | |
| | | | KΖ, | LC, | LK, | LR, | LS, | LT, | LU, | LV, | MD, | MG, | MK, | MN, | MW, | MX, | NO, | NZ, | |
| | | | PL, | PT, | RO, | RU, | SD, | SE, | SG, | SI, | SK, | SL, | TJ, | TM, | TR, | TT, | UA, | UG, | |
| | | | UZ, | VN, | YU, | ZW | | | | | | | | | | | | | |
| | | RW: | GH, | KE. | LS. | MW. | SD, | SZ, | UG, | ZW. | AT. | BE, | CH, | DE, | DK, | ES, | FI, | FR, | |
| | | | | | | | | | | | | BF, | | | | | | | |
| | | | | | | | | TD, | | | | | | | | . , | | . , | |
| | US | 6207 | | | | | | | | | US : | 1996- | 7151 | 63 | | 1 | 9960 | 917 | < |
| | CA | 2265 | 828 | | | | | | | | | 1997- | | | | | 9970 | | |
| | | | | | | | | | | | | 1997- | | | | | 9970 | 917 | < |
| | | 7435 | | | | | | | | | | | | - | | _ | | | |
| | | | | | | | | | | | ED . | 1997- | 9452 | 49 | | 1 | 9970 | 917 | / |
| | | | | | | | | | | | | IT, | | | | | | | |
| | | 14. | | FI. | | DL, | DIC, | шо, | 111, | GD, | OI. | , 11, | шт, | шо, | иш, | υ . , | nic, | , | |
| | TD | 2001 | | | | т | | 2001 | 0221 | | TD : | 1998- | 51/0 | 0.4 | | 1 | 0070 | 017 | |
| | | | | | | | | | | | | 2002- | | | | | | | |
| PRIO | | | | | | n | | 2002 | 0323 | | | 1996- | | | | | | | \ |
| PRIOR | KIII | APP | LIN. | TMFO | . : | | | | | | | 1996-
1995- | | | | | | | |
| | | | | | | | | | | | | | | | | | | | |
| | | | | | | | | | | | | 1995- | | | | | 9950 | | |
| | | | | | | | | | | | | 1996- | | | | | | | |
| | | | _ | | | | | | | | | 1997- | | | | | | | |
| | | | | | | | | | | | | duncin. | | | | | | | |

Materials and methods are provided for producing patterned multi-array, multi-sp. surfaces for use in diagnostics. The invention provides for electrochemiluminescence methods for detecting or measuring an analyte of interest. It also provides for novel electrodes for ECL assays. Materials and methods are provided for the chemical and/or phys. control of conducting domains and reagent deposition for use multiply specific

testing procedures. REFERENCE COUNT: THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS

L42 ANSWER 28 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1997:341994 CAPLUS

127:34643

DOCUMENT NUMBER:

TITLE: Polymers of N-acrylovlmorpholine derivative activated at one end and conjugates with bioactive materials and

surfaces

INVENTOR(S): Veronese, Francesco M.; Schiavon, Oddone; Caliceti,

Paolo; Sartore, Luciana; Ranucci, Elisabetta; Ferruti, Paolo

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

PATENT ASSIGNEE(S): Consiglio Nazionale Delle Ricerche, Italy

SOURCE: U.S., 9 pp. CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------|------|----------|-----------------|------------|
| | | | | |
| US 5629384 | A | 19970513 | US 1994-243869 | 19940517 < |

19950607 <--US 5631322 A 19970520 US 1995-475177 19950607 FO.: US 1994-243869 A3 19940517 PRIORITY APPLN. INFO.:

AB The title polymers having a single reactive moiety at one end of the polymer chain have the following structure R-Z-X-Y (R = N-acryloylmorpholine residue with d.p. 6-280, which yields number-average mol.

weight 1000-40,000; Z-X-Y = polymer capping moiety; X = saturated residue of linear or branched aliphatic series CrH2r, r = 1-12; Y = reactive moiety, such as -OH, -CO2H, or -NH2; Z = moiety that readily reacts to cap a polymer free radical, e.g., S). The monofunctional polymer is a suitable alternative to monofunctional PEG for modification of substances having biol. and biotech. applications.

L42 ANSWER 29 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1997:192127 CAPLUS

DOCUMENT NUMBER: 126 • 185989

TITLE: Preparation of (aminostyryl)pyridinium compounds for

radiolabelling cell membranes

INVENTOR(S): Lambert, Carol; Mease, Ronnie C.; Mcafee, John G. Research Corporation Technologies, Inc., USA; Lambert, PATENT ASSIGNEE(S): Carol; Mease, Ronnie C.; Mcafee, John G.

PCT Int. Appl., 38 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent.

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE -----______ WO 9702246 A1 19970123 WO 1995-US8460 19950706 <--W: CA, JP, MX

RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE A1 19970123 CA 1995-2225861 19950706 <--WO 1995-US8460 W 19950706 CA 2225861 PRIORITY APPLN. INFO.:

OTHER SOURCE(S): MARPAT 126:185989

AB (E)-R1ZCH:CHC6H4(NR2)-4 X (Z = pyridinio-4-yl)[I; R = CnH2n+1; R1 = organic group containing detectable radioisotope (sic); X = biol. acceptable anion; n = 4-16] were prepared Thus, R2CH:CHC6H4(NR2)-4 (R = decv1, R2 = 4-pyridv1) was N-alkylated by (E)-Bu3SnCH:CHCH2OTs (preparation given) and the product iodinated to give I [R = decyl, R1 = (E)-125ICH:CHCH2, X unspecified]. Data for biol. activity of I were given.

L42 ANSWER 30 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1997:169157 CAPLUS DOCUMENT NUMBER: 126:225315

TITLE: Bicyclic heterocyclic derivatives having

al-adrenergic and 5HT1A serotonergic activities Leonardi, Amedeo; Motta, Gianni; Riva, Carlo; Testa, INVENTOR(S):

Rodolfo

Recordati S.A., Chemical and Pharmaceutical Company, PATENT ASSIGNEE(S): Switz.

SOURCE: U.S., 84 pp., Cont.-in-part of U.S. 5,474,994.

CODEN: USXXAM DOCUMENT TYPE: Pat.ent.

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3 PATENT INFORMATION:

> PATENT NO. KIND DATE APPLICATION NO. DATE

| US | 5605896 | A | 19970225 | US | 1994-299188 | | 19940831 | < |
|----------|---------------|--------|------------|----|-------------|----|----------|---|
| US | 5403842 | A | 19950404 | US | 1992-888775 | | 19920526 | < |
| AU | 9336296 | A | 19930913 | AU | 1993-36296 | | 19930223 | < |
| RO | 112111 | В3 | 19970530 | RO | 1994-1404 | | 19930223 | < |
| PL | 175556 | B1 | 19990129 | PL | 1993-304889 | | 19930223 | < |
| RU | 2128656 | C1 | 19990410 | RU | 1994-43324 | | 19930223 | < |
| SK | 280143 | B6 | 19990910 | SK | 1994-1007 | | 19930223 | < |
| ZA | 9301278 | A | 19931118 | ZA | 1993-1278 | | 19930224 | < |
| LT | 3038 | В | 19940925 | LT | 1993-354 | | 19930224 | < |
| CN | 1079738 | A | 19931222 | CN | 1993-105852 | | 19930526 | < |
| CN | 1040434 | В | 19981028 | | | | | |
| US | 5474994 | A | 19951212 | US | 1993-67861 | | 19930526 | < |
| FI | 9403876 | A | 19940823 | FI | 1994-3876 | | 19940823 | < |
| NO | 9403140 | A | 19940825 | NO | 1994-3140 | | 19940825 | < |
| PRIORITY | APPLN. INFO.: | | | IT | 1992-MI408 | Α | 19920225 | |
| | | | | US | 1992-888775 | A2 | 19920526 | |
| | | | | US | 1993-67861 | A2 | 19930526 | |
| | | | | EP | 1993-301264 | A | 19930222 | |
| | | | | WO | 1993-EP420 | A | 19930223 | |
| OTHER SC | DURCE (S): | MARPAT | 126:225315 | | | | | |

OTHER SOURCE(S): MARPAT 126:225315 GI

AB Bicyclic heterocyclic derivs, such as I [X = N, O, S; W = C(O), C(S), CH(OH), bond; R2 = H, optionally substituted alkyl, alkenyl, alkylnyl, carbocycle, heterocycle; R3 = alkyl, hydroxyalkyl, Ph, OH, alkoxy, alkoxyalkyl; R6 = H, halogen, NO2, NH2, AcNH, mono-, dialkylamino, CN, OH, alkoxy, alkyl; Y = CO, CO2, CONH, CH(OH), CH:CH, CH:CHCO2, CH:CHCONH, CH2NH, CH2NHCO, CH2NHSO2, CH2S, NH, NHCO, NHCONH, NHSO2, O, S, SOZHH, CONHO, CSNH, NHCO2, COS, CONH(CH2)m, m = 1-6; Z = N, A = (un) substituted Ph, pyrimidinyl, 1, 4-benzodioxan-8-vl, benzopyran-8-vl,

benzofuran-7-yl, dihydrobenzopyran-8-yl; Z = CH2N; Z = CH, A = one or two Ph, 4-FC6H4CO, 2-oxo-1-benzimidazolinyl, (CH2)nOA, n = 0-2], and their pharmaceutically acceptable salts useful as $\alpha 1$ -adrenergic and 5HT1A serotonergic agents for the treatment of hypertension, urethral and lower urinary tract contractions, and other disorders are described. Thus, benzopyran II was prepared by heating 1-(2-methoxyphenyl)piperazine with benzopyran III at 180° for 5 h. II had IC50 = 29 nM for $\alpha 1$ -adrenergic receptor binding, IC50 = 9 nM for 5HT1A receptor binding, ED25 = 45 μ g/kg i.v. hypotensive effect and ED25 = 1.4 μ g/kg in Na-induced urethral contractility assays.

L42 ANSWER 31 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1996:452240 CAPLUS

ACCESSION NUMBER: 1996:45224 DOCUMENT NUMBER: 125:221638

TITLE: Nonpeptidal P2 Ligands for HIV Protease Inhibitors: Structure-Based Design, Synthesis, and Biological

Evaluation

AUTHOR(S): Ghosh, Arun K.; Kincaid, John F.; Walters, D. Eric; Chen, Yan; Chaudhuri, Narayan C.; Thompson, Wayne J.; Culberson, Chris; Fitzgerald, Paula M. D.; Lee, Hee Yoon; et al.

CORPORATE SOURCE: Department of Chemistry, University of Illinois,

Chicago, IL, 60607, USA
SOURCE: Journal of Medicinal Chemistry (1996), 39(17),

3278-3290 CODEN: JMCMAR; ISSN: 0022-2623

CODEN: JMCMAR; ISSN: 0022-2623 American Chemical Society Journal

English

PUBLISHER: DOCUMENT TYPE: LANGUAGE: GI

AB Design and synthesis of nonpeptidal bis-tetrahydrofuran ligands based upon the X-ray crystal structure of the HIV-I protease-inhibitor Ro 31-8959 led to replacement of two amide bonds and a l0r-aromatic system of Ro 31-8959 class of HIV protease inhibitors. Detailed structure-activity studies have now established that the position of ring oxygens, ring size, and stereochem. are all crucial to potency. Of particular interest, I with (38, 385, 683)-bis-Thf is the most potent inhibitor (ICSO value 1.8 ± 0.2 nM; CICSO value 46 ± 4 nM) in this series. The X-ray structure of protein-inhibitor I has provided insight into the ligand-binding site interactions. As it turned out, both oxygens in the bis-Thf ligands are involved in hydrogen-bonding interactions with Asp 29 and Asp 30 NH present in the S2 subsite of HIV-I protease. Stereoselective routes have been developed to obtain these novel ligands in optically pure form

L42 ANSWER 32 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1996:35000 CAPLUS

DOCUMENT NUMBER: 124:232248

TITLE: Benzopyran derivatives having affinity for

al-adrenergic and 5HT1A-serotoninergic receptors INVENTOR(S): Leonardi, Amedeo; Motta, Gianni; Riva, Carlo; Testa,

Rodolfo PATENT ASSIGNEE(S): Recordati S.A., Chemical and Pharmaceutical Company,

Switz. U.S., 37 pp. Cont.-in-part of U.S. 5,403,842.

SOURCE:

CODEN: USXXAM DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | | DATE |
|------------------------|--------|-----------|-----------------------|--------|---------------|
| US 5474994 | A | 19951212 | US 1993-67861 | | 19930526 < |
| US 5403842 | A | 19950404 | US 1992-888775 | | 19920526 < |
| EP 558245 | A1 | 19930901 | EP 1993-301264 | | 19930222 < |
| R: AT, BE, CH, | DE, DK | , ES, FR, | GB, GR, IE, IT, LI, I | LU, MO | C, NL, PT, SE |
| AU 9336296 | A | 19930913 | AU 1993-36296 | | 19930223 < |
| RO 112111 | B3 | 19970530 | RO 1994-1404 | | 19930223 < |
| PL 175556 | B1 | 19990129 | PL 1993-304889 | | 19930223 < |
| SK 280143 | B6 | 19990910 | SK 1994-1007 | | 19930223 < |
| CN 1079738 | A | 19931222 | CN 1993-105852 | | 19930526 < |
| CN 1040434 | В | 19981028 | | | |
| FI 9403876 | A | 19940823 | FI 1994-3876 | | 19940823 < |
| NO 9403140 | A | 19940825 | NO 1994-3140 | | 19940825 < |
| US 5605896 | A | 19970225 | US 1994-299188 | | 19940831 < |
| PRIORITY APPLN. INFO.: | | | US 1992-888775 | A2 | 19920526 |
| | | | EP 1993-301264 | A | 19930222 |
| | | | IT 1992-MI408 | A | 19920225 |
| | | | WO 1993-EP420 | A | 19930223 |
| | | | US 1993-67861 | A2 | 19930526 |
| OTHER SOURCE(S): | MARPAT | 124:23224 | .8 | | |

OTHER SOURCE(S): MARPAT 124:232248

GI

This invention provides bicyclic heterocyclic derivs. I wherein the dotted line represents a single or double bond; X represents a nitrogen, oxygen or sulfur atom, or an amino or alkylamino group, a sulfinyl or sulfonyl group; W represents a carbonyl, thiocarbonyl, hydroxymethylene, or a methylene group or a bond; or when X is nitrogen and W is a methine, the fused rings represent a quinoline; R2 represents, e.g, a hydrogen atom or an alkyl, alkenyl, alkynyl, carbocyclic or heterocyclic group, each of which groups may optionally be substituted; or R2 itself represents a trifluoromethyl or an aroyl group; R3 represents a hydrogen atom or an alkyl, hydroxyalkyl, alkyl-O-R4 Ph, hydroxy, or O-R4, wherein R4 represents an alkyl group optionally substituted with an aryl group; R6 represents a hydrogen or halogen atom or a nitro, amino, acylamino, alkylsulfonylamino, alkylamino, dialkylamino, cyano, hydroxy, alkoxy or alkyl group; R7 represents a hydrogen atom or an alkoxy group; Y = e.g., CO, COO, CONH; Z represents a linear or branched chain alkylene group having from 1 to 6 carbon atoms and optionally having one hydroxy substituent; B = e.g., II, n = 1 or 2, A = substituted Ph, 2-pyrimidinyl; and their pharmaceutically acceptable salts useful for the treatment of hypertension, urethral and lower urinary tract contractions, and other disorders. The compds. are also useful for binding α1-adrenergic and 5HT1A serotonergic receptors, in vitro or in vivo. Thus, e.g., esterification of 8-carboxy-3-methyl-4-oxo-2-phenyl-4H-1-benzopyran with 1-(3-chloropropyl)-4-(2-methoxyphenyl)piperazine followed by HCl treatment afforded 8-{3-[4-(2-methoxyphenyl)-1-piperazinyl]propoxycarbonyl}-3-methyl-4-oxo-2-phenyl-4H-1-benzopyran dihydrochloride (III.2HCl) which exhibited IC50's of 20 and 19 nM, resp., for al and 5-HT1A receptor binding. Data were also presented for the effect of I on K+ stimulation of rat bladder strips, and on urethral contractions and blood pressure in dogs.

III

L42 ANSWER 33 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1995:996262 CAPLUS

DOCUMENT NUMBER: 124:56360

preparation of new retinol (vitamin A) derivatives and their use in pharmaceuticals and cosmetics

INVENTOR(S): Weischer, Carl Heinrich; Oestreich, Wolfgang

PATENT ASSIGNEE(S): Germany

TITLE:

STN Search - 10/517,692

SOURCE: Ger. Offen., 9 pp.
CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

DE 4415204 A1 19951102 DE 1994-4415204 19940430 <-PRIORITY APPLN. INFO.: DE 1994-4415204 19940430

OTHER SOURCE(S): CASREACT 124:56360

GT

AB Preparation of retinol esters (I; R = H, Ac) of salicylic and acetylsalicylic acids and their use in pharmaceuticals and cosmetics are described. Pharmaceutical applications claimed include antiinflammatories, geriatric disorders, dermatol., cytoprotectives, antineuralgics, antitumor agents, antithrombotics, antidepenerative action. As therapeutics and cosmetics, I can be used for prophylaxis or treatment of inflammation and other skin and appendage disorders, such as, sunburn, vesicular pityriasis, dandruff, seborrhea, eczema, and pyoderma of the scalp, seborrheic eczema of the hair bed, seborrheic companion symptoms of androgenetic alopecia and other skin diseases including neurodermitis, psoriasis, hyperkeratosia, urticaria, and of the hair follicles. I can be used in therapy for night blindness, necrosia, intoxication, tumor sicknesses (e.g. bronchial carcinoma), neuralgia, old age problems, vitamin A deficiency diseases, and as prophylaxis for thrombosis.

L42 ANSWER 34 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1994:708324 CAPLUS

DOCUMENT NUMBER: 121:308324

TITLE: Antibody-drug conjugates for parenteral administration
INVENTOR(S): Barton, Russell Lawern; Guttman-Carlisle, Deborah

Lane; Koppel, Gary Allen

PATENT ASSIGNEE(S): Eli Lilly and Co., USA
SOURCE: Eur. Pat. Appl., 22 pp.

CODEN: EPXXDW DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PA | TENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|----|----------------|--------|-------------|-----------------------|------------|
| | | | | | |
| EP | 620011 | A1 | 19941019 | EP 1994-302059 | 19940322 < |
| | R: AT, BE, CH, | DE, DK | ES, FR, GB, | , GR, IE, IT, LI, LU, | NL, PT, SE |
| US | 5556623 | A | 19960917 | US 1993-40323 | 19930330 < |

| CA 211 | .9733 | A1 | 19941001 | CA | 1994-2119733 | | 19940322 | < |
|-------------|--------------|----|----------|----|--------------|----|----------|---|
| JP 063 | 321880 | A | 19941122 | JΡ | 1994-57065 | | 19940328 | < |
| US 564 | 13573 | A | 19970701 | US | 1995-541847 | | 19951010 | < |
| US 566 | 5358 | A | 19970909 | US | 1996-649568 | | 19960517 | < |
| PRIORITY AP | PPLN. INFO.: | | | US | 1993-40323 | A | 19930330 | |
| | | | | US | 1995-541847 | A3 | 19951010 | |
| | | | | | | | | |

OTHER SOURCE(S): MARPAT 121:308324

Immunoconjugates of antibodies or antigen-recognizing fragments of antibodies and monovalent cytotoxic drug derivs. make use of β-alanine derived linkers, wherein the antibody or fragment thereof is attached to the linker's carboxy group via an ester or amide group and the drug is attached through the linker's 2-position methylene group. Intermediates, compns. and methods of use also are provided. For example, MeCOC(:CHOEt)CONHCH2CH2CO2H was prepared and reacted with desacetylvinblastine hydrazide sulfate, then with antibody VX 007B to give a conjugate. An anticancer activity of the conjugate was tested in vivo against xenografts of UCLA/P3 lung adenocarcinoma in mice.

L42 ANSWER 35 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1994:676197 CAPLUS

DOCUMENT NUMBER: 121:276197

TITLE: Thin-film hydrophilic polar multi-functionalized

polymer (HPMP) matrix systems and methods for constructing and displaying ligands

INVENTOR(S): Hudson, Derek; Johnson, Charles R.; Ross, Michael J.;

Shoemaker, Kevin R.; Cass, Robert T.; Giebel, Lutz B.; Zhou, Peng

PATENT ASSIGNEE(S):

Arris Pharmaceutical Corp., USA SOURCE: PCT Int. Appl., 59 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

| PA | TENT NO. | | | KIN | D DAT | Œ | AP | PLICAT | ION NO. | | DATE | |
|---------|----------|-------|-----|-----|--------|-------|-------|--------|---------|-----|-------------|---|
| | | | | | | | | | | | | |
| WO | 9419694 | | | A1 | 199 | 40901 | WO | 1994- | US2036 | | 19940218 | < |
| | W: AU | , CA, | CN, | JP, | NO, US | i | | | | | | |
| | RW: Al | , BE, | CH, | DE, | DK, ES | , FR, | GB, G | R, IE, | IT, LU, | MC, | NL, PT, SE | |
| US | 5576220 | | | A | 199 | 61119 | US | 1993- | 19725 | | 19930219 | < |
| US | 5585275 | | | A | | | | | | | 19930618 | |
| WO | 9405394 | | | A1 | 199 | 40317 | WO | 1993- | US8267 | | 19930902 | < |
| | W: AU | , CA, | JP, | NO | | | | | | | | |
| | RW: Al | | | | | | | | | | NL, PT, SE | |
| | 9463939 | | | | | | | | | | 19940218 | |
| JP | 0850760 | 2 | | T | 199 | 60813 | JP | 1994- | 519285 | | 19940218 | < |
| PRIORIT | Y APPLN. | INFO | . : | | | | US | 1993- | 19725 | - 2 | A 19930219 | |
| | | | | | | | US | 1993- | 79741 | - 2 | A 19930618 | |
| | | | | | | | WO | 1993- | US8267 | | A 19930902 | |
| | | | | | | | US | 1992- | 939065 | | A2 19920902 | |
| | | | | | | | WO | 1994- | US2036 | 1 | W 19940218 | |

AB Methods and systems of unhindered construction and display of tethered organic ligand mols. are disclosed, especially preparation and use of thin film.

substantially non-crosslinked hydrophilic polar multi-functionalized polymers (HPMPs) anchored to a variety of functionalized substrates so that the HPMP forms a thin film matrix layer providing a unique, highly hydrated, high dielec. environment equivalent to an aqueous solution, for affinity

binding of ligands to tagged target mols. Preparation of e.g. a HPMP (dextranized) polyethylene substrate surface is described, as is e.g. production of dextrans containing masking functional groups. In a test of stability to TFA of the product of the invention as compared to a epichlorohydrin-bonded dextran-polyethylene, after only 2 h, approx. 90% of dextrans were lost from the epichlorohydrin-bonded surfaces, whereas only minor loss (<10%) was detected from the surface stapled according to the invention. Use of the The HPMP for peptide synthesis and peptide library screening is also described.

L42 ANSWER 36 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1993:625779 CAPLUS

DOCUMENT NUMBER: 119:225779

TITLE: Design and synthesis of novel ligands for the 5-HT3

and the 5-HT4 receptor

AUTHOR(S): Blum, E.; Buchheit, K. H.; Buescher, H. H.; Gamse, R.; Kloeppner, E.; Meigel, H.; Papageorgiou, C.; Waelchli,

R.; Revesz, L. Preclin. Res., Sandoz Pharma AG, Basel, CH-4002,

CORPORATE SOURCE: Switz.

SOURCE: Bioorganic & Medicinal

Chemistry Letters (1992),

2(5), 461-6

CODEN: BMCLE8; ISSN: 0960-894X

DOCUMENT TYPE: Journal LANGUAGE:

English

OTHER SOURCE(S): CASREACT 119:225779

GI

A novel highly potent 5-HT3 antagonist and Tropisetron analog I is described with an increased efficacy to inhibit cisplatin induced emesis in ferrets. Four novel structural classes of gastroprokinetic benzamide bioisosteres, e.g., II, are presented. 5-HT derivs., e.g., III, are described as ligands of the recently discovered 5-HT4 receptor.

ACCESSION NUMBER: 1992:123815 CAPLUS

DOCUMENT NUMBER: 116:123815

A polyclonal antibody preparation with Michaelian TITLE:

catalytic properties

Gallacher, Gerard; Jackson, Caroline S.; Searcey, AUTHOR(S):

Mark; Badman, Geoffrey T.; Goel, Rajiv; Topham, Christopher M.; Mellor, Geoffrev W.; Brocklehurst,

CORPORATE SOURCE: Queen Mary and Westfield Coll., Univ. London, London,

El 4NS, UK

SOURCE: Biochemical Journal (1991), 279(3), 871-81

CODEN: BIJOAK; ISSN: 0306-3275

DOCUMENT TYPE: Journal

LANGUAGE: English

4-Nitrophenyl 4'-(3-aza-2-oxoheptyl)phenyl carbonate (I), an amide conjugate (II) involving the carboxy group of 4-nitrophenyl

4'-carboxymethylphenyl phosphate and an amino group of keyhole-limpet hemocyanin, and a fluorescein derivative (III) were synthesized. II was used as an immunogen with which to raise polyclonal antibodies in

multigeneration cross-bred sheep; III was used for the initial assessment of the antisera via binding assays monitored by fluorescence polarization; I was used as a chromogenic substrate for the investigation of catalytic activity. The IgG from the antiserum of sheep number 270 was isolated by

Na2SO4 precipitation and chromatog. on Protein G-Sepharose. This preparation of IqG

catalyzed the hydrolysis of I; the catalysis at pH 8.0 and 25° obeyed Michaelis-Menten kinetics with at least 25 turnovers, Km = 3.34 μM, and lower limits for kcat of 0.029 s-1 and for kcat/Km of 8.77 × 103 M-1-s-1, on the unlikely assumption that the concentration of catalytic antibody is provided by twice the total IgG concentration (two sites per mol.); probable ests. of the fraction of the total IgG that is anti-haptenic IgG and of the fraction of this that is catalytically active suggest that the values of kcat/Km are actually very much larger than these lower limits. The failure of the antibody preparation to catalyze the hydrolysis of the isomeric 2-nitrophenyl carbonate (IV) which differs from I only in the position of the nitro substituent in the leaving group, compels the view that catalytic activity is due to antibody rather than contaminant enzyme; this conclusion is supported by (a) the failure of the following to discriminate effectively between the isomeric substrates I and IV: pig liver carboxylesterase, rabbit liver carboxylesterase (collectively EC 3.1.1.1), whole serum from a non-immunized sheep and whole serum from a sheep immunized with a derivative of 3-0methylnoradrenaline and (b) the lack of catalytic activity in IgG prepns. from sheep immunized with sulfoxide or sulfone analogs of immunogen II. The various parameters used for the comparison of the kinetic characteristics of hydrolytic catalytic antibodies are discussed. The characteristics of hydrolysis of I catalyzed by the present polyclonal antibody preparation are shown to be substantially better in most respects than those of analogous reactions of two other carbonate esters catalyzed by monoclonal antibodies.

L42 ANSWER 38 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1989:115144 CAPLUS

DOCUMENT NUMBER: 110:115144

TITLE: Derivatives of all-trans- and 13-cis-retinoic acid and

their preparation

INVENTOR(S): Deluca, Hector F.; Kutner, Andrzej; Schnoes, Heinrich

к.

PATENT ASSIGNEE(S): Wisconsin Alumni Research Foundation, USA

SOURCE: PCT Int. Appl., 13 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. ----WO 8707604 19871217 WO 1987-US1276 A1 19870601 <--W: CH, DE, GB, JP RW: AT, BE, CH, DE, FR, GB, IT, LU, NL, SE A US 4757140 19880712 US 1986-869791 19860602 <--EP 271552 A1 19880622 EP 1987-904165 19870601 <--B1 19931027 EP 271552 R: AT, BE, CH, DE, FR, GB, IT, LI, LU, NL, SE JP 01500190 T 19890126 JP 1987-503792 19870601 <--JP 06051716 В 19940706 AT 96432 Т 19931115 AT 1987-904165 19870601 <--CA 1305136 C 19920714 CA 1987-538880 19870604 <--A 19890620 A 19901030 US 4841038 US 1988-190443 19880505 <--19890323 <--US 4966965 US 1989-327540 A 19860602 A 19870601 W 19870601 PRIORITY APPLN. INFO.: US 1986-869791 EP 1987-904165 WO 1987-US1276 W 19870601 US 1988-190443 A3 19880505 OTHER SOURCE(S): CASREACT 110:115144

AB all-trans- And 13-cis-retinoic acid (I; R = CO2H, R1 = H; R = H, R1 = CO2H, resp.) derivs, are prepared all-trans-I (R = CO2H, R1 = H) (III) was stirred with an equimolar mixture of N-hydroxysuccinimide and DCC in dioxane at room temperature to give III succinimido ester, which was treated with a solution of CoA in THF at pH 8.0-8.5 at 35° under N to give III CoA ester.

L42 ANSWER 39 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

1988:406321 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 109:6321

Preparation of haloalkylazetidinones TITLE:

INVENTOR(S): Miller, Marvin Joseph University of Notre Dame, USA Eur. Pat. Appl., 13 pp. PATENT ASSIGNEE(S):

SOURCE:

CODEN: EPXXDW DOCUMENT TYPE: Pat.ent.

English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

> PATENT NO. KIND DATE APPLICATION NO. DATE

| EP 2 | 256763 | | A1 | 1 | 9880 | 224 | | ΕP | 1987-306916 | | 19870805 | < |
|-----------|----------|---------|------|-------|------|------|-----|-----|---------------|---|----------|---|
| EP 2 | 256763 | | B1 | 1 | 9931 | 208 | | | | | | |
| | R: AT, | BE, CH, | DE, | FR, | GB, | IT, | LI, | NI | , SE | | | |
| US 4 | 1751296 | | A | 1 | 9880 | 614 | | US | 1986-893748 | | 19860806 | < |
| IL 8 | 33420 | | A | 1 | 9920 | 715 | | IL | 1987-83420 | | 19870803 | < |
| JP 6 | 3044560 | | A | 1 | 9880 | 225 | | JΡ | 1987-196152 | | 19870805 | < |
| JP (| 7080839 | | В | 1 | 9950 | 830 | | | | | | |
| HU 4 | 15011 | | A2 | 1 | 9880 | 530 | | HU | 1987-3580 | | 19870805 | < |
| HU 2 | 201737 | | В | 1 | 9901 | 228 | | | | | | |
| CA I | 1282066 | | С | 1 | 9910 | 326 | | CA | 1987-543734 | | 19870805 | < |
| AT 9 | 8229 | | T | 1 | 9931 | 215 | | ΑT | 1987-306916 | | 19870805 | < |
| PRIORITY | APPLN. 3 | INFO.: | | | | | | US | 1986-893748 | A | 19860806 | |
| | | | | | | | | ΕP | 1987-306916 | A | 19870805 | |
| OTHER SOU | JRCE(S): | | CASE | REACT | 109 | :632 | 21; | MAF | RPAT 109:6321 | | | |
| GI | | | | | | | | | | | | |

The title compds. [I; R = protected NH2, alkyl, phenylalkyl; R1 = alkyl, alkoxy, (un) substituted Ph, PhO, PhCH2O; R2 = H, alkyl, CH:CHR3, (CH2)mCHO, (CH2)nOR4 (CH2)p X1, (CH2)qCO2R5; R3 = H, alkyl, CO2R5, Ph, alkoxyphenyl, furyl; R4 = hydroxy protective group; R5 = carboxy protective group; X, X1 = C1, Br, iodo; m, n, p, q = 0-2] were prepared by reaction of R2CH:CHCHRCONHOCOR1 with a weak base in the presence of a pos. halogen reagent. MeSOCH2CH2CH(NHCO2CH2Ph)CO2Me (preparation given) was stirred vigorously at 180-190° for 1.5-2 h to give, after ester hydrolysis, H2C:CHCH(NHCO2CH2Ph)CO2H which was esterified with N-hydroxysuccinimide and the product amidated with HONH2-HCl. The N-hydroxyamide thus obtained was condensed with ClCO2CH2Ph to give H2C:CHCH(NHCO2CH2Ph)CONHOCO2CH2Ph which, in MeCN, was stirred with K2CO3 followed by addition of H2O and then Br to give azetidinone II.

L42 ANSWER 40 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1988:22078 CAPLUS

DOCUMENT NUMBER: 108:22078

TITLE: Synthesis of coenzyme A ester of retinoic acid:

intermediate in vitamin A metabolism

Kutner, Andrzei; Renstrom, Britta; Schnoes, Heinrich AUTHOR(S):

K.; DeLuca, Hector F.

Coll. Agric. Life Sci., Univ. Wisconsin, Madison, WI, CORPORATE SOURCE:

53706, USA

Proceedings of the National Academy of Sciences of the SOURCE:

United States of America (1986), 83(18), 6781-4

CODEN: PNASA6; ISSN: 0027-8424

Journal

DOCUMENT TYPE: LANGUAGE: English

AB CoA esters of all-trans-I (R = CO2H; Rl = H) and 13-cis-retinoic acid I (R = H; Rl = CO2H) were prepared for use in studying vitamin A metabolism, from I (R,Rl = H, CO2H) via their activated succinimido esters or anhydrides.

L42 ANSWER 41 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1987:598057 CAPLUS

DOCUMENT NUMBER: 107:198057

TITLE: Synthesis and immobilization of a novel acridine

derivative on microparticulate silica. A study of its interactions with single-stranded oligonucleotides by

high-performance liquid chromatography

AUTHOR(S): Bischoff, Rainer; Regnier, Fred E.

CORPORATE SOURCE: Dep. Biochem., Purdue Univ., West Lafayette, IN, 47907, USA

SOURCE: Journal of Chromatography (1987), 397, 13-24 CODEN: JOCRAM: ISSN: 0021-9673

DOCUMENT TYPE: Journal

LANGUAGE: English

AB A novel approach for immobilizing acridine on 5-µm silica gel is

described. The acridine moiety is functionalized with a carboxylic acid group at its reactive 9-position and activated, leading to

9-acridinylpropionic acid N-hydroxysuccinimide ester. This derivative is efficiently bound to the silica matrix through a primary aliphatic amine group at the end of a 15-atom spacer arm. The chromatog, properties of the final stationary phases, as evaluated with d(T)10 and d(A)10 at various pH values and organic solvent conces., resemble those of hydrophobic weak anion exchangers. When a secondary amine group is placed closed to

the acridine moiety in 1 of the packings, enhanced binding of the oligodeoxyribonucleotides is observed that goes beyond a purely additive effect.

L42 ANSWER 42 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1987:67117 CAPLUS

DOCUMENT NUMBER: 106:67117

TITLE: Compounds for site-enhanced delivery of radionuclides

and their uses

INVENTOR(S): Bodor, Nicholas S.

PATENT ASSIGNEE(S): University of Florida, USA SOURCE: PCT Int. Appl., 262 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|----------------|---------|----------|-----------------|------------|
| | | | | |
| WO 8600898 | A1 | 19860213 | WO 1985-US1334 | 19850715 < |
| W: AU. DK. FT. | NO. IIS | | | |

RW: AT, BE, CH, DE, FR, GB, IT, LU, NL, SE

| AU | 8546358 | | A | 19860225 | AU 1985-46358 | | 19850715 | < |
|----------|----------|--------|--------|-------------|----------------|----|----------|---|
| EP | 187832 | | A1 | 19860723 | EP 1985-903633 | | 19850715 | < |
| | R: AT, | BE, C | H, DE, | FR, GB, IT, | LI, LU, NL, SE | | | |
| JP | 61106556 | | A | 19860524 | JP 1985-160040 | | 19850719 | < |
| ZA | 8505476 | | A | 19870325 | ZA 1985-5476 | | 19850719 | < |
| CA | 1267899 | | A1 | 19900417 | CA 1985-487165 | | 19850719 | < |
| ES | 552072 | | A1 | 19870416 | ES 1986-552072 | | 19860217 | < |
| NO | 8600981 | | A | 19860520 | NO 1986-981 | | 19860314 | < |
| FI | 8601118 | | A | 19860318 | FI 1986-1118 | | 19860318 | < |
| DK | 8601247 | | A | 19860520 | DK 1986-1247 | | 19860318 | < |
| US | 4963688 | | A | 19901016 | US 1987-88523 | | 19870821 | < |
| US | 5155227 | | A | 19921013 | US 1990-561920 | | 19900802 | < |
| PRIORITY | APPLN. | INFO.: | | | US 1984-632314 | A2 | 19840719 | |
| | | | | | WO 1985-US1334 | A | 19850715 | |
| | | | | | US 1986-879120 | B1 | 19860319 | |
| | | | | | US 1987-88523 | A3 | 19870821 | |

OTHER SOURCE(S): MARPAT 106:67117

GI For diagram(s), see printed CA Issue.
AB A composition of matter comprised: (1

A composition of matter comprised: (1) the residue of a chelating agent having ≥1 reactive functional group selected from NH2, CO2H, OH, amide, and imide, said functional group being not essential for the complexing properties of the chelating agent, said residue being characterized by the absence of a H atom from ≥1 of said reactive functional groups of said chelating agent which is either (a) capable of chelating with a metallic radionuclide or (b) chelated with a metallic radionuclide; and (2) a dihydropyridine/pyridinium salt redox carrier moiety; said chelating agent residue and said carrier moiety being coupled to each other to form a hydrolytically cleavable linkage between them. More specifically, a salt I (A = residue of chelating agent capable of chelating with a metal radionuclide; y = 1,2; [QC+] is the hydrophilic, ionic pyridinium salt form of a dihydropyridine/pyridinium salt redox carrier; X- = anion of a pharmaceutically acceptable organic or inorg. acid; n = valence of acid anion; m = number which when multiplied by n = y. This complex provides a new radionuclide pharmaceutical that, in its lipoidal dihydropyridine form, penetrates the blood-brain barrier and allows increased levels of radionuclide concentration in the brain. This radionuclide delivery system is well suited for use in scintigraphy and similar radiog. techniques. Homocysteine thiolactone II in THF reacted with (H2NCH2)2 to give H2NCH2CH2NHCOCH(CH2CH2SH)NHCOCH2N(CO2CMe3)CH2CH2SH (III). Esterification of nicotinic acid with N-hydroxysuccinimide gave the succinimidyl ester, which was quaternized with MeI to give succinimidyl trigonellinate (IV). Amidation of IV with III gave homocysteinamide V (R = CO2CMe3), deblocking of which with HCl(q) in EtOH gave V (R = H). This in EtOH containing NaOH was treated with 99mTcO4- and Na2S2O4 solution to give the complex between dihydropyridine VI and the oxotechnetate-99m ion.

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L42 ANSWER 43 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                        1984:570730 CAPLUS
DOCUMENT NUMBER:
                        101:170730
ORIGINAL REFERENCE NO.: 101:25811a, 25814a
TITLE:
                        Nitroaliphatic compounds and their use
INVENTOR(S):
                        Okamoto, Masanori; Iwami, Morita; Takase, Shigehiro;
                        Uchida, Itsuo; Umehara, Kazuyoshi; Kohsaka, Masanobu;
                         Imanaka, Hiroshi
PATENT ASSIGNEE(S):
                        Fujisawa Pharmaceutical Co., Ltd., Japan
SOURCE:
                         Eur. Pat. Appl., 58 pp.
                        CODEN: EPXXDW
DOCUMENT TYPE:
                        Patent
LANGHAGE .
                        English
FAMILY ACC. NUM. COUNT: 1
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PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPI | LICATION NO. | | DATE | |
|--|---------|--------------|------|--------------|----|----------|---|
| | | | | | - | | |
| EP 113106 | A1 | 19840711 | EP : | 1983-112955 | | 19831222 | < |
| EP 113106 | B1 | 19860514 | | | | | |
| R: AT, BE, CH, | DE, FR | , GB, IT, LI | , LU | , NL, SE | | | |
| ZA 8308831 | A | 19840725 | ZA : | 1983-8831 | | 19831125 | < |
| AU 8321744 | A | 19840705 | AU : | 1983-21744 | | 19831128 | < |
| AU 560980 | A
B2 | 19870430 | | | | | |
| US 4767768 | A | 19880830 | US : | 1983-559260 | | 19831208 | < |
| CA 1231949 | A1 | 19880126 | CA : | 1983-443153 | | 19831213 | < |
| FI 8304702
FI 78904
FI 78904
AT 19773 | A | 19840701 | FI : | 1983-4702 | | 19831221 | < |
| FI 78904 | В | 19890630 | | | | | |
| FI 78904 | C | 19891010 | | | | | |
| AT 19773 | | 19860515 | AT : | 1983-112955 | | 19831222 | < |
| JP 59152366 | | 19840831 | JP : | 1983-252520 | | 19831227 | < |
| JP 02019822 | В | 19900507 | | | | | |
| DK 8306077 | A | 19840701 | | 1983-6077 | | 19831230 | < |
| NO 8304884 | A | 19840702 | NO : | 1983-4884 | | 19831230 | < |
| NO 158379 | В | 19880524 | | | | | |
| NO 158379 | С | 19880831 | | | | | |
| HU 32795 | A2 | 19840928 | HU : | 1983-4543 | | 19831230 | < |
| HU 200747 | В | 19900828 | | | | | |
| ES 528561 | | 19850501 | | 1983-528561 | | 19831230 | |
| SU 1389678 | A3 | 19880415 | | 1983-3678551 | | 19831230 | |
| US 4778804 | A | 19881018 | | 1985-786754 | | 19851011 | |
| US 4782088 | A | 19881101 | | 1986-946868 | | 19861229 | |
| US 4863926 | | 19890905 | | 1987-119091 | | 19871110 | |
| JP 02160750 | | 19900620 | JP : | 1989-259680 | | 19891004 | < |
| JP 04017944 | В | 19920326 | | | | | |
| PRIORITY APPLN. INFO.: | | | | | | 19821231 | |
| | | | | 1983-559260 | | 19831208 | |
| | | | | 1983-112955 | | 19831222 | |
| | | | US : | 1985-786754 | A3 | 19851011 | |
| | | | | | | | |

OTHER SOURCE(S): CASREACT 101:170730

Mitrated oximes RCR1(NO2)CR2:CR3C(:NOR4)R5 and RCR1(NO2)CHR2CHR3C(:NOR4)R5 (R = H, alky1, alkoxypheny1; R1 = H, alky1; R2 = alky1; R3 = H, alky1; R4 = H, alky1, carboxyalky1, carbalkoxyalky1; R5 = H, CH:NOH, cyano, a N-piperazinecarbonyl group, alkanoyl, esterified CO2H, alky1, CONH2, substituted carbamoyl), which were prepared, showed antiplatelet aggregation and vasodilator activity. Thus, (E,E)-MeCH:CEtCH:CHCONH2 was treated with NaNO2 at pH 3.0 to give (E)-MeCH(NO2)CEt:CHC(:NOH)CONH2, which also exhibited antihypertensive activity.

L42 ANSWER 44 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1984:45461 CAPLUS
DOCUMENT NUMBER: 100:45461
DDTCINAL DEFERENCE NO. 100:45663

ORIGINAL REFERENCE NO.: 100:6863a,6866a

TITLE: A new enkephalin analog: trans-4-hydroxycinnamoylqlvcvl-qlvcvl-phenvlalanvl-leucine. Synthesis and

biological properties

AUTHOR(S): Amar, Claudine; Vilkas, Erna; Laurent, Stephane;

Gautray, Bruno; Schmitt, Henri
CORPORATE SOURCE: Lab. Org. Biol. Chem., Univ. Paris-Sud, Orsay, Fr.

SOURCE: International Journal of Peptide & Protein Research

(1983), 22(4), 434-6

CODEN: IJPPC3; ISSN: 0367-8377

DOCUMENT TYPE: Journal

LANGUAGE: English

AB A leucine-enkephalin [58822-25-6] analog in which the N-terminal tyrosine has been replaced by trans-4-hydroxycinnamic acid was synthesized by

liquid-phase coupling methods. The central cardiovascular effects of this analog were investigated and the results discussed.

L42 ANSWER 45 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1982:582827 CAPLUS
DOCUMENT NUMBER: 97:182827

DOCUMENT NUMBER: 9/:18282/

ORIGINAL REFERENCE NO.: 97:30608h,30609a

TITLE: Grafting of cysteine and cystine to the surface of an aerosil across an amide bond

AUTHOR(S): Filippov, A. P.; Kozynchenko, A. P.

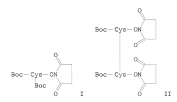
CORPORATE SOURCE: Inst. Fiz. Khim. im. Pisarzhevskogo, Kiev, USSR

SOURCE: Ukrainskii Khimicheskii Zhurnal (Russian Edition)

(1982), 48(8), 860-3 CODEN: UKZHAU; ISSN: 0041-6045

DOCUMENT TYPE: Journal

LANGUAGE: Russian



AB Cysteine I (Boc = Me3CO2C) and cystine II were grafted onto an aerosil by treating with aminoaerosil RSiMe2NH2.

L42 ANSWER 46 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1980:198783 CAPLUS DOCUMENT NUMBER: 92:198783

ORIGINAL REFERENCE NO.: 92:32226h,32227a

TITLE: Glucosamine peptide derivatives and their

pharmaceutical compositions
INVENTOR(S): Yuichi, Yamamura; Ichiro, Azuma; Shigeru, Kobayashi

PATENT ASSIGNEE(S): Takeda Chemical Industries, Ltd., Japan

SOURCE: Eur. Pat. Appl., 80 pp.

CODEN: EPXXDW DOCUMENT TYPE: Patent

LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------|------|----------|-----------------|------------|
| | | | | |
| EP 2677 | A1 | 19790711 | EP 1978-101524 | 19781202 < |
| EP 2677 | B1 | 19821013 | | |

| R: | CH, DE, | FR, GB, I | T | | | | |
|----------------|----------|-----------|--------------|----------------|----|----------|---|
| JP 54079 | 227 | A | 19790625 | JP 1977-145415 | | 19771202 | < |
| JP 54079 | 228 | A | 19790625 | JP 1977-145416 | | 19771202 | < |
| JP 02033 | 719 | В | 19900730 | | | | |
| JP 54120 | 696 | A | 19790919 | JP 1978-28012 | | 19780310 | < |
| JP 63000 | 446 | В | 19880107 | | | | |
| US 44302 | 65 | A | 19840207 | US 1982-393870 | | 19820630 | < |
| PRIORITY APPL | N. INFO. | . : | | JP 1977-145415 | | 19771202 | |
| | | | | JP 1977-145416 | | 19771202 | |
| | | | | JP 1978-28012 | | 19780310 | |
| | | | | US 1978-962033 | A1 | 19781120 | |
| | | | | US 1981-249902 | A1 | 19810401 | |
| OTHER SOURCE (| S): | MARPA | AT 92:198783 | | | | |

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB Acetylmuramyl dipeptide derivs. I [n = 0, R = H, alkyl; n = 1-9, R = H, NH2; R1 and R2 = alkyl; R3 and R4 = (R2OH, R5 and R6 = CO2H, CONH2; R7 = H, R8CC (R8 = acyclic hydrocarbon which can be o-substituted by cycloalkyl), Q (1 = 1-9; m = 0-9; t = 2-100; R8 and R9 = H, alkyl; R10 = alkyl, CO2H which can be esterified, OH which can be etherified, pyrrolidino which can be substituted)] were prepared as immunostimulants. Thus, acetylmuramyl dipeptide II (R11 = H) was esterified with Z-B-Ala-CO6H4NO2-p (Z = PHCH2OZC) to give II (R11 = Z-B-Ala), which was hydrogenated over Pd/C to give B-alanylmuramic acid derivative III (R12 = H) (IV). IV was N-acylated with CR2:CMSCOSU (Su = succinimido) to give III (R12 = CH2:CMSCO) (V), which was polymerized to give the homopolymer of V. V was copolymd. with N-vinyl-2-pyrrolidone, stearyl vinyl ether, and tridecyl methacrylate to give the resp. copolymers. The cell-mediated immunostimulatory activities of several I were tested.

L42 ANSWER 47 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1979:474325 CAPLUS DOCUMENT NUMBER: 91:74325

ORIGINAL REFERENCE NO.: 91:12008h,12009a

TITLE: Alkylaniline compounds and an antiatherosclerosis

agent containing it American Cyanamid Co., USA

PATENT ASSIGNEE(S): American Cyanamid Co., U SOURCE: Ger. Offen., 263 pp. CODEN: GWXXBX.

DOCUMENT TYPE: Patent

LANGUAGE: German

FAMILY ACC. NUM. COUNT: 5 PATENT INFORMATION:

| PA: | TENT NO. | KIND | DATE | API | PLICATION NO. | DATE | |
|-----|----------|------|----------|-----|---------------|----------|---|
| | | | | | | | |
| DE | 2841707 | A1 | 19790427 | DE | 1978-2841707 | 19780925 | < |
| US | 4117158 | A | 19780926 | US | 1977-836946 | 19770927 | < |
| ZA | 7805071 | A | 19791128 | ZA | 1978-5071 | 19780906 | < |
| BE | 870687 | A1 | 19791128 | BE | 1978-190650 | 19780922 | < |
| US | 4254138 | A | 19810303 | US | 1979-87137 | 19791022 | < |
| US | 4272546 | A | 19810609 | US | 1979-87136 | 19791022 | < |
| DK | 7904815 | A | 19800516 | DK | 1979-4815 | 19791114 | < |
| SE | 7909398 | A | 19800516 | SE | 1979-9398 | 19791114 | < |

| ES 485942 | A1 | 19801101 | ES 1979-485942 | 19791114 < |
|------------------------|----|----------|----------------|-------------|
| US 4309553 | A | 19820105 | US 1980-156144 | 19800603 < |
| PRIORITY APPLN. INFO.: | | | US 1977-836946 | A 19770927 |
| | | | US 1977-836947 | A 19770927 |
| | | | US 1977-861736 | A3 19771219 |
| | | | GB 1978-44562 | A 19781115 |

AB More than 100 4-RRINKCH4R2 (I; R = CB-19-alkyl; R1 = H or a group convertible in vivo to H, e.g., Me, MeCO, CH2SC3Ma; R2 = alkoxycarbonyl, substituted carbamoyl or carboximidoyl, alkoxyalkyl, acyl, cyanoalkyl, etc.), useful in the treatment or prevention of atherosclerosis (no data), were prepared Thus, 4-O2NC6H4SO2NH2 was treated with NaH, and the 4-O2NC6H4SO2NHNa treated with 4-Me(CH2)15NHC6H4COC1 (prepared from the acid) to give I (R = H, R1 = hexadecyl, R2 = 4-O2NC6H4SO2NH).

L42 ANSWER 48 OF 48 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1978:89892 CAPLUS

DOCUMENT NUMBER: 88:89892

ORIGINAL REFERENCE NO.: 88:14095a,14098a

TITLE: All-trans-retinoic acid esters and amides

Ι

INVENTOR(S): Gander, R. J.; Gurney, J. A.

KIND DAT

PATENT ASSIGNEE(S): Johnson and Johnson, USA

SOURCE: Belg., 20 pp. CODEN: BEXXAL

DOCUMENT TYPE: Patent LANGUAGE: French

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO.

PRIORITY APPLN. INFO.:

BE 847942 197 A1 US 4108880 Α 197 CA 1062700 197 A1 NL 7612201 Α 197 GB 1543824 Α 197 US 4190594 198

| Έ | API | PLICATION NO. | | DATE | |
|--------|-----|---------------|---|----------|---|
| | | | - | | |
| 770503 | BE | 1976-172046 | | 19761103 | < |
| 80822 | US | 1975-628177 | | 19751103 | < |
| 90918 | CA | 1976-264672 | | 19761102 | < |
| 770505 | NL | 1976-12201 | | 19761103 | < |
| 90411 | GB | 1976-45746 | | 19761103 | < |
| 800226 | US | 1978-906168 | | 19780515 | < |
| | US | 1975-628177 | Α | 19751103 | |
| | | | | | |

AB All-trans-retinoic acid derivs. I (R = 2-cyclohexylethoxy, MeO2C(CH2)100, BO(CE2)40, cholesteryl, 3-CH2:CHG6H4CH20, 4-ERCEH4CH20, 4-BrC6H4CH20, OCH2COR1, NHPr, NHCMe3, NHCMe2CH2CMe3, morpholino, 4-HOC6H4NH, 4,2-MeO2C(HO)C6H3NH, 3,4-(MeO)2C6H3CH2CH2WH, 2-benzothiazolyl,lamino, 1-imidazolyl, 2-nicotinoylhydrazino, 1-benzotriazolyl, 1,2,4-triazol-1-yl, B-ionone hydrazono N-cyclohexylaminocarbonyl-N-cyclohexylamino; Rl = cholesteryloxy, Ph, 4-BrC6H4, 4-MeC6H4, 4-PhC6H4, 2,5-(MeO)2C6H3, 3,4-ChC0)2C6H3, 3,4-(MeO)3C6H2, 2,4,6-MeO3C6H2, 2,4,

were prepared for use as sunscreen agents (no data). Thus K all-trans-retinoate was treated with Br(CH2)10CO2Me to give I [R = O(CH2)10CO2Me].

| => log h
COST IN U.S. DOLLARS | SINCE FILE
ENTRY | TOTAL |
|--|---------------------|------------------|
| FULL ESTIMATED COST | 169.52 | 267.62 |
| DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) | SINCE FILE
ENTRY | TOTAL
SESSION |
| CA SUBSCRIBER PRICE | -38.40 | -44.80 |

SESSION WILL BE HELD FOR 120 MINUTES
STN INTERNATIONAL SESSION SUSPENDED AT 09:30:07 ON 31 JAN 2008

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PASSWORD:

** * * * * RECONNECTED TO STN INTERNATIONAL * * * * * * * SESSION RESUMED IN FILE 'CAPLUS' AT 09:48:27 ON 31 JAN 2008 FILE 'CAPLUS' ENTERED AT 09:48:27 ON 31 JAN 2008 COPYRIGHT (C) 2008 AMERICAN CHEMICAL SOCIETY (ACS)

| ### SESSION 169.52 267.62 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE ENTRY SESSION - 44.80 => fil caplus | COST IN U.S. DOLLARS | SINCE FILE | TOTAL |
|--|--|------------|---------|
| DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION -38.40 -44.80 | | ENTRY | SESSION |
| CA SUBSCRIBER PRICE -38.40 C-44.80 -> fil caplus COST IN U.S. DOLLARS SINCE FILE ENTRY SESSION FULL ESTIMATED COST 169.52 267.62 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE ENTRY SESSION ENTRY SESSION ENTRY SESSION ENTRY SESSION ENTRY SESSION | FULL ESTIMATED COST | 169.52 | 267.62 |
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| FULL ESTIMATED COST 169.52 267.62 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION | | | |
| ENTRY SESSION | FULL ESTIMATED COST | | |
| CA SUBSCRIBER PRICE -38.40 -44.80 | DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) | | |
| | CA SUBSCRIBER PRICE | -38.40 | -44.80 |

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FILE COVERS 1907 - 31 Jan 2008 VOL 148 ISS 5 FILE LAST UPDATED: 30 Jan 2008 (20080130/ED)

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=>

Uploading C:\Program Files\Stnexp\Queries\10527694-31Jan08.str

chain nodes :

CHEAT HOUSE: 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37 38 39 40 41 42 43 49 50 51 57 58 59 60 61 62 63 ring nodes : 44 45 46 47 48 52 53 54 55 56 chain bonds : $1-2 \quad 2-3 \quad 2-8 \quad 3-4 \quad 4-5 \quad 4-7 \quad 5-6 \quad 5-9 \quad 10-11 \quad 11-12 \quad 11-16 \quad 12-13 \quad 12-17 \quad 13-14$ 14-15 14-18 19-20 20-21 20-30 21-22 22-23 22-26 22-28 23-24 24-25 24-31 26-27 26-29 32-33 33-34 33-41 34-35 35-36 35-39 36-37 37-38 37-43 39-40 39-44 94 55-51 48-50 52-59 53-58 54-60 56-57 60-61 60-62 60-63

```
ring bonds :
44-45 44-48 45-46 46-47 47-48 52-53 52-56 53-54 54-55 55-56
exact/norm bonds :
4-7 12-17 22-28 44-45 44-48 44-49 45-46 45-51 46-47 47-48 48-50 52-53
52-56 52-59 53-54 53-58 54-55 54-60 55-56 56-57 60-61 60-62 60-63
exact bonds :
2-3 3-4 4-5 11-12 12-13 13-14 20-21 21-22 22-23 22-26 23-24 33-34 34-35
35-36 35-39 36-37
normalized bonds :
1-2 2-8 5-6 5-9 10-11 11-16 14-15 14-18 19-20 20-30 24-25 24-31 26-27
26-29 32-33 33-41 37-38 37-43 39-40 39-42
Match level :
1:CLASS 2:CLASS 3:CLASS 4:CLASS 5:CLASS 6:CLASS 7:CLASS 8:CLASS 9:CLASS
10:CLASS 11:CLASS 12:CLASS 13:CLASS 14:CLASS 15:CLASS 16:CLASS 17:CLASS
18:CLASS 19:CLASS 20:CLASS 21:CLASS 22:CLASS 23:CLASS 24:CLASS 25:CLASS
26:CLASS 27:CLASS 28:CLASS 29:CLASS 30:CLASS 31:CLASS 32:CLASS 33:CLASS
34:CLASS 35:CLASS 36:CLASS 37:CLASS 38:CLASS 39:CLASS 40:CLASS 41:CLASS
42:CLASS 43:CLASS 44:Atom 45:Atom 46:Atom 47:Atom 48:Atom 49:CLASS 50:CLASS 51:CLASS 52:Atom 53:Atom 54:Atom 55:Atom 56:Atom 57:CLASS 58:CLASS
59:CLASS 60:CLASS 61:CLASS 62:CLASS 63:CLASS
```

L43 STRUCTURE UPLOADED

=> d

L43 HAS NO ANSWERS

L43 STR

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

Structure attributes must be viewed using STN Express query preparation.

=> s 143

REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress...
Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

SAMPLE SEARCH INITIATED 09:49:10 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 1 TO ITERATE

100.0% PROCESSED 1 ITERATIONS 0 ANSWERS SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**

PROJECTED ITERATIONS: 1 TO 80
PROJECTED ANSWERS: 0 TO 0

L44 0 SEA SSS SAM L43

L45 0 L44

=> log h

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ENTRY SESSION
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PASSWORD:

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COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 0.48 269.04 DISCOUNT AMOUNTS (FOR OUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE 0.00 -44.80 => fil caplus COST IN U.S. DOLLARS SINCE FILE TOTAL. ENTRY SESSION FULL ESTIMATED COST 0.48 269.04 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE 0.00 -44.80

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546575 "SULFATE"

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FILE LAST UPDATED: 30 Jan 2008 (20080130/ED)
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=> s crosslinking agent
        209268 CROSSLINKING
           136 CROSSLINKINGS
        209328 CROSSLINKING
                 (CROSSLINKING OR CROSSLINKINGS)
       884233 AGENT
       1302570 AGENTS
       1822121 AGENT
                (AGENT OR AGENTS)
L46
        72274 CROSSLINKING AGENT
                 (CROSSLINKING(W) AGENT)
=> s sulfate
        546575 SULFATE
         99691 SULFATES
        595653 SULFATE
L47
                 (SULFATE OR SULFATES)
=> s 146 and 147
         2066 L46 AND L47
L48
=> s chrondroitin sulfate
           10 CHRONDROITIN
        546575 SULFATE
        99691 SULFATES
        595653 SULFATE
                 (SULFATE OR SULFATES)
L49
             8 CHRONDROITIN SULFATE
                 (CHRONDROITIN(W)SULFATE)
=> s (chondroitin sulfate OR "Chondroitin, hydrogen sulfate")
         16310 CHONDROITIN
           100 CHONDROITINS
         16325 CHONDROITIN
                 (CHONDROITIN OR CHONDROITINS)
        546575 SULFATE
        99691 SULFATES
        595653 SULFATE
                 (SULFATE OR SULFATES)
         13463 CHONDROITIN SULFATE
                 (CHONDROITIN(W)SULFATE)
         16310 "CHONDROITIN"
           100 "CHONDROITINS"
         16325 "CHONDROITIN"
                 ("CHONDROITIN" OR "CHONDROITINS")
       1039920 "HYDROGEN"
          6143 "HYDROGENS"
       1043336 "HYDROGEN"
                 ("HYDROGEN" OR "HYDROGENS")
```

99691 "SULFATES" 595653 "SULFATE"

("SULFATE" OR "SULFATES")

52 "CHONDROITIN, HYDROGEN SULFATE"

("CHONDROITIN" (W) "HYDROGEN" (W) "SULFATE") L50 13474 (CHONDROITIN SULFATE OR "CHONDROITIN, HYDROGEN SULFATE")

=> s 146 and 150

116 L46 AND L50 L51

=> s 151 and biomaterial

10076 BIOMATERIAL 10856 BIOMATERIALS

16264 BIOMATERIAL

(BIOMATERIAL OR BIOMATERIALS) L52 8 L51 AND BIOMATERIAL

=> d ibib abs 1-8

L52 ANSWER 1 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

2006:234728 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 144:299306

TITLE: Process for isolating biomaterial from tissue and an isolated biomaterial extract prepared therefrom

INVENTOR(S): Ying, Jackie Y.; Pek, Shona

PATENT ASSIGNEE(S): Agency for Science, Technology and Research, Singapore

SOURCE: PCT Int. Appl., 20 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

| PAT | TENT | NO. | | | KIN | D | | | | | | ION 1 | | | | ATE | |
|------|-------|------|------|-----|-----|-----|------|------|-----|------|-------|-------|-----|-----|-----|------|-----|
| WO | 2006 | 0284 | 15 | | A1 | _ | | | | | | | | | | | |
| | W: | ΑE, | AG, | AL, | AM, | AT, | AU, | AZ, | BA, | BB, | BG, | BR, | BW, | BY, | BZ, | CA, | CH, |
| | | CN, | CO, | CR, | CU, | CZ, | DE, | DK, | DM, | DZ, | EC, | EE, | EG, | ES, | FI, | GB, | GD, |
| | | GE, | GH, | GM, | HR, | HU, | ID, | IL, | IN, | IS, | JP, | KE, | KG, | KP, | KR, | KZ, | LC, |
| | | LK, | LR, | LS, | LT, | LU, | LV, | MA, | MD, | MG, | MK, | MN, | MW, | MX, | MZ, | NA, | NI, |
| | | NO, | NZ, | OM, | PG, | PH, | PL, | PT, | RO, | RU, | SC, | SD, | SE, | SG, | SK, | SL, | SY, |
| | | TJ, | TM, | TN, | TR, | TT, | TZ, | UA, | UG, | US, | UZ, | VC, | VN, | YU, | ZA, | ZM, | ZW |
| | RW: | AT, | BE, | BG, | CH, | CY, | CZ, | DE, | DK, | EE, | ES, | FI, | FR, | GB, | GR, | HU, | IE, |
| | | ΙT, | LU, | MC, | NL, | PL, | PT, | RO, | SE, | SI, | SK, | TR, | BF, | ВJ, | CF, | CG, | CI, |
| | | CM, | GA, | GN, | GQ, | GW, | ML, | MR, | ΝE, | SN, | TD, | TG, | BW, | GH, | GM, | KE, | LS, |
| | | MW, | MZ, | NA, | SD, | SL, | SZ, | TZ, | UG, | ZM, | ZW, | AM, | ΑZ, | BY, | KG, | KZ, | MD, |
| | | RU, | TJ, | TM | | | | | | | | | | | | | |
| ΑU | 2004 | 3230 | 01 | | A1 | | 2006 | 0316 | | AU 2 | 004- | 3230 | 01 | | 2 | 0040 | 909 |
| EP | 1786 | 829 | | | A1 | | 2007 | 0523 | | EP 2 | 004- | 7756 | 11 | | 2 | 0040 | 909 |
| | | DE, | | | | | | | | | | | | | | | |
| RITY | Y APP | LN. | INFO | . : | | | | | | WO 2 | 004 - | SG28: | 9 | | A 2 | 0040 | 909 |

PRIO A process for isolating a biomaterial extract from tissue is disclosed.

The process comprises the step of contacting the tissue with an extracting solution so as to extract a biomaterial into solution A solution containing the

biomaterial extract is separated before being freeze-dried at a rate sufficient to enable the biomaterial to be isolated. The examples relate to the extraction of collagen from skin or hide using an acetic acid solution as the solvent. The product obtained may be used in cosmetic, medical, pharmaceutical, food, or veterinarian industries.

REFERENCE COUNT: 8 THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD, ALL CITATIONS AVAILABLE IN THE RE FORMAT

L52 ANSWER 2 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2005:346799 CAPLUS

DOCUMENT NUMBER: 142:397837

TITLE: Protein biomaterials and biocoacervate

INVENTOR(S): Masters, David B.; Berg, Eric P.

PATENT ASSIGNEE(S): Gel-Del Technologies, Inc., USA SOURCE: PCT Int. Appl., 99 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| | DATE |
|--|-------------|
| WO 2005034852 A2 20050421 WO 2004-US27975
WO 2005034852 A3 20071213 | 20040826 |
| W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, | BZ, CA, CH, |
| CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, | FI, GB, GD, |
| GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, | |
| LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, | MZ, NA, NI, |
| NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, | SK, SL, SY, |
| TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, | |
| RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, | ZM, ZW, AM, |
| AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, | CZ, DE, DK, |
| EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, | PT, RO, SE, |
| SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, | ML, MR, NE, |
| SN, TD, TG, AP, EA, EP, OA | |
| AU 2004279349 A1 20050421 AU 2004-279349 | 20040826 |
| CA 2537315 A1 20050421 CA 2004-2537315 | |
| US 2006073207 A1 20060406 US 2004-929117 | |
| EP 1660013 A2 20060531 EP 2004-782454 | 20040826 |
| R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, | SE, MC, PT, |
| IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, | |
| PRIORITY APPLN. INFO.: US 2003-497824P | P 20030826 |
| WO 2004-US27975 | W 20040826 |

The present invention relates to protein biocoacervates and biomaterials and the methods of making and using protein biocoacervates and biomaterials. More specifically the present invention relates to protein biocoacervates and biomaterials that may be utilized for various medical applications including, but not limited to, drug delivery devices for the controlled release of pharmacol, active agents, coated medical devices (e.g., stents, valves), vessels, tubular grafts, vascular grafts, wound healing devices including protein suture biomaterials and biomeshes, dental plugs and implants, skin/bone/tissue grafts, tissue fillers, protein biomaterial adhesion prevention barriers, cell scaffolding and other biocompatible biocoacervate or biomaterial devices. Soluble bovine collagen was dissolved in water. To this solution was added elastin and sodium heparinate dissolved in water. The

elastin/heparin solution was added quickly to the collagen solution with minimal

stirring thereby immediately producing an amorphous coacervate precipitate

L52 ANSWER 3 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2004:857457 CAPLUS

DOCUMENT NUMBER: 141:337851

TITLE: Molded elastin article and process for producing the

AUTHOR(S):

```
same
INVENTOR(S):
                            Miyamoto, Keiichi; Kitazono, Eiichi; Miyoshi,
                            Takanori: Kaneko, Hiroaki: Sumi, Yoshihiko: Hirata,
                            Hitoshi
PATENT ASSIGNEE (S):
                            Teijin Limited, Japan
SOURCE:
                            PCT Int. Appl., 22 pp.
                            CODEN: PIXXD2
DOCUMENT TYPE:
                            Patent
LANGUAGE:
                            Japanese
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
     PATENT NO.
                           KIND DATE
                                                APPLICATION NO.
                                                                           DATE
                           ----
                                                 ______
     WO 2004087232
                           A1 20041014 WO 2004-JP4494
                                                                           20040330
          W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,
              CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
              GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI,
          LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, RW, BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN,
              TD, TG
     AU 2004226551
                             A1
                                    20041014 AU 2004-226551
                                                                            20040330
     CA 2520704
                             A1
                                    20041014 CA 2004-2520704
                                                                            20040330
     EP 1609492
                             A1
                                    20051228 EP 2004-724354
                                                                            20040330
          R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
              IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK
     CN 1767864
                                   20060503
                                                  CN 2004-80008680 20040330
                             A
     US 2006194036
                             A1
                                    20060831
                                                  US 2005-551545
                                                                            20050930
PRIORITY APPLN. INFO.:
                                                  JP 2003-94398
                                                                       A 20030331
                                                  WO 2004-JP4494
                                                                       A 20040330
OTHER SOURCE(S):
                           MARPAT 141:337851
    Disclosed is a molded elastin article in which a fiber structure made of
     aliphatic polyester fibers having an average fiber diameter of 0.05 to 50 µm
are
     employed as a supporting base and which is flexible, bioabsorbable and has
     such tear strength as allowing stitching in practice. This molded elastin
     article is useful as a material for tubes and artificial vessels to be
     used in transplantation in vivo which are bioabsorbable and have such tear
     strength and flexibility as withstanding stitching during surgery
     operations. A tube made with polylactic acid (Lacty 9031) was reacted
     with elastin and a water-soluble crosslinking agent prepared from
     dodecanediarboxylic acid and 4-hydroxyphenyldimethyl sulfonium
     methylsulfate to obtain a elastin-crosslinked polyester tube.
                                   THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                            5
                                   RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L52 ANSWER 4 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                            2003:493623 CAPLUS
DOCUMENT NUMBER:
                            140:169587
TITLE:
                            Preparation and evaluation of molecularly-defined
                            collagen-elastin-glycosaminoglycan scaffolds for
```

tissue engineering

Kuppevelt, T. H.

Daamen, W. F.; van Moerkerk, H. Th. B.; Hafmans, T.;

Buttafoco, L.; Poot, A. A.; Veerkamp, J. H.; van

CORPORATE SOURCE: NCMLS, Department of Biochemistry 194, University Medical Centre Nijmegen, Nijmegen, 6500 HB, Neth.

SOURCE: Biomaterials (2003), 24(22), 4001-4009

CODEN: BIMADU; ISSN: 0142-9612

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

AB Extracellular matrix components are valuable building blocks for the preparation of biomaterials involved in tissue engineering, especially if their biol., chemical and phys. characteristics can be controlled. In this study, isolated type I collagen fibrils, elastin fibers and chondroitin sulfate (CS) were used for the preparation of molecularly-defined collagen-elastin-glycosaminoglycan scaffolds. A total of 12 different scaffolds were prepared with four different ratios of collagen and elastin (1:9, 1:1, 9:1 and 1:0), with and without chemical crosslinking, and with and without CS. Collagen was essential to fabricate coherent, porous scaffolds. Electron microscopy showed that collagen and elastin phys. interacted with each other and that elastin fibers were enveloped by collagen. By carbodiimide-crosslinking, amine groups were coupled to carboxvlic groups and CS could be incorporated. More CS could be bound to collagen scaffolds (10%) than to collagen-elastin scaffolds (2.4-8.5% depending on the ratio). The attachment of CS increased the water-binding capacity to up to 65%. Scaffolds with a higher collagen content had a higher tensile strength whereas addition of elastin increased elasticity. Scaffolds were cytocompatible as was established using human myoblast and fibroblast culture systems. It is concluded that molecularly-defined

composite scaffolds can be composed from individual, purified, extracellular matrix components. Data are important in the design and application of tailor-made biomaterials for tissue engineering. REFERENCE COUNT: 26 THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L52 ANSWER 5 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1997:503015 CAPLUS

DOCUMENT NUMBER: 127:113411

TITLE: Use of injectable or implantable biomaterials for filling or blocking lumens and voids of the body

INVENTOR(S): Rhee, Woonza M.; Berg, Richard A.; Chu, George H.;
Delustro, Frank A.; Jolivette, Dan M.; Mccullough,

Kimberly A.

PATENT ASSIGNEE(S): Collagen Corporation, USA

SOURCE: PCT Int. Appl., 28 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

KIND DATE APPLICATION NO. DATE PATENT NO. A1 19970626 WO 1996-US20553 WO 9722372 19961218 W: AU, CA, JP RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE CA 2239772 A1 19970626 CA 1996-2239772 19961218 AU 9713473 A 19970714 AU 1997-13473 19961218 AU 708320 B2 19990729
EP 876166 A1 19981111 EP 1996-945006
EP 876166 B1 20040818 19961218 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,

| JP 2000501975 | T | 20000222 | JP | 1997-523044 | | 19961218 |
|------------------------|----|----------|----|--------------|----|----------|
| AT 273722 | T | 20040915 | AT | 1996-945006 | | 19961218 |
| ES 2227627 | Т3 | 20050401 | ES | 1996-945006 | | 19961218 |
| JP 2005320352 | A | 20051117 | JP | 2005-211983 | | 20050721 |
| PRIORITY APPLN. INFO.: | | | US | 1995-574050 | A | 19951218 |
| | | | JP | 1997-523044 | A3 | 19961218 |
| | | | WO | 1996-US20553 | W | 19961218 |

AB Methods for completely or partially blocking, augmenting, sealing, or filing various biol. lumens and voids within the body of a patient are disclosed. Lumens include arteries, veins, intestines, fallopian tubes, and trachea. Voids include various lesions, fissures, diverticulae, cysts, fistulae, aneurysms, or other undesirable voids that may exist within a patient's body. An effective amount of a biomaterial is administered (e.g., via injection, catheter, or surgical implantation) into the lumen or void. Thus, fibrillar collagen (55 mg/mL) was mixed with PEG succinimidyl glutarate (SG-PEG) in a 1-10 molar ratio of collagen-SG-PEG. The collagen/SG-PEG reaction mixture was extruded into small diameter tubings. The above collagen rod was inserted into each of the ureters of a guinea pig cadaver and cut to size. The crosslinked collagen rod was not dislodged and the bladder did not leak, as viewed under UV light.

L52 ANSWER 6 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1996:660894 CAPLUS

DOCUMENT NUMBER: 125:285011

TITLE: Use of hydrophobic crosslinking agents to prepare

crosslinked biomaterial implants

INVENTOR(S): Rhee, Woonza M.

PATENT ASSIGNEE(S): Collagen Corporation, USA

SOURCE: Eur. Pat. Appl., 26 pp. CODEN: EPXXDW

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|--------|-------------|-----------------|-------------|
| EP 732109 | A1 | 19960918 | EP 1996-102366 | 19960216 |
| R: AT, CH, DE, | FR, GB | , IT, LI, N | L, SE | |
| CA 2165728 | A1 | 19960915 | CA 1995-2165728 | 19951220 |
| JP 09249751 | A | 19970922 | JP 1996-58138 | 19960314 |
| US 6962979 | B1 | 20051108 | US 1999-344230 | 19990625 |
| US 2004121951 | A1 | 20040624 | US 2003-448246 | 20030528 |
| US 7129209 | B2 | 20061031 | | |
| US 2005154125 | A1 | 20050714 | US 2004-997246 | 20041123 |
| JP 2006181389 | A | 20060713 | JP 2006-66762 | 20060310 |
| PRIORITY APPLN. INFO.: | | | US 1995-403358 | A 19950314 |
| | | | JP 1996-58138 | A3 19960314 |
| | | | US 1997-987467 | B1 19971209 |
| | | | US 1999-344230 | A1 19990625 |

AB Novel crosslinked biomaterial compns. are prepared using hydrophobic polymers as a crosslinking agent. Preferred hydrophobic polymers are those that contain two or more reactive succinimidyl groups, including disuccinimidyl suberate (I), bis(sulfosuccinimidyl suberate, and dithiobis(succinimidyl propionate). Crosslinked biomaterial compns. prepared using mixts. of hydrophobic and hydrophilic crosslinking agents are also disclosed. The compns. of the present invention can be used to prepare formed implants for use in a variety of medical applications. Thus, 1.0 mt of 35 mg/ml collapen was mixed with 3 mg I in a syringe and

incubated at 37° for 16 h. The crosslinked collagen material was extruded out of the plunger end of the syringe and the resulting crosslinked cylindrical gels were then sectioned into 5 mm thick disks. The solubilization of crosslinked collagen in trypsin solution and oxidative degradation in 3% H2O2 was 7 and 14 days, resp. After implantation of the crosslinked collagen in rats for 90 days it had a discrete, football-shaped, bolus-like configuration, whereas noncrosslinked formulation was present as a more diffuse mass.

L52 ANSWER 7 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1996:599239 CAPLUS

DOCUMENT NUMBER: 125:285010 TITLE: Method of preparing crosslinked polymeric

biomaterial compositions for use in tissue

augmentation

INVENTOR(S): Rhee, Woonza M.; Berg, Richard A.; Rosenblatt, Joel S.; Tefft, Jacqueline A.; Braga, Larry J.; Smestad,

Thomas L. PATENT ASSIGNEE(S): USA

PATENT NO. KIND DATE

SOURCE: U.S., 14 pp., Cont.-in-part of U.S. Ser. No. 236,769.

APPLICATION NO.

DATE

A 19950630

CODEN: USXXAM DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 18 PATENT INFORMATION:

| | FAIENI NO. | KIND | DATE | AFFILICATION NO. | DAIL |
|------|--------------------|------|----------|------------------|-------------|
| | US 5550187 | A | 19960827 | US 1994-287549 | 19940808 |
| | US 5162430 | A | 19921110 | US 1989-433441 | 19891114 |
| | US 5328955 | A | 19940712 | US 1992-922541 | 19920730 |
| | US 5304595 | A | 19940419 | US 1992-998802 | 19921230 |
| | US 5306500 | A | 19940426 | US 1993-110577 | 19930823 |
| | US 5376375 | A | 19941227 | US 1994-177578 | 19940105 |
| | US 5413791 | A | 19950509 | US 1994-198128 | 19940217 |
| | US 5475052 | A | 19951212 | US 1994-236769 | 19940502 |
| | US 5523348 | A | 19960604 | US 1994-292415 | 19940818 |
| | US 5543441 | A | 19960806 | US 1995-427576 | 19950424 |
| | US 5527856 | A | 19960618 | US 1995-440274 | 19950512 |
| | US 5643464 | A | 19970701 | US 1995-497573 | 19950630 |
| | EP 697218 | A2 | 19960221 | EP 1995-112218 | 19950803 |
| | EP 697218 | A3 | 19960529 | | |
| | R: DE, FR, GB, | IT | | | |
| PRIO | RITY APPLN. INFO.: | | | US 1988-274071 | B2 19881121 |
| | | | | US 1989-433441 | A2 19891114 |
| | | | | US 1992-922541 | A3 19920730 |
| | | | | US 1994-198128 | A2 19940217 |
| | | | | US 1994-236769 | A2 19940502 |
| | | | | US 1992-930142 | A3 19920814 |
| | | | | US 1993-110577 | A3 19930823 |
| | | | | US 1994-177578 | A3 19940105 |
| | | | | US 1994-287549 | A3 19940808 |
| | | | | US 1994-292415 | A3 19940818 |
| | | | | | |

US 1995-497573 AB The present invention discloses a novel method for preparing crosslinked biomaterial compns. for use in the augmentation of soft or hard tissue. In general, the method comprises mixing a biocompatible polymer, which is preferably collagen, with a sterile, dry crosslinking agent, which is preferably a synthetic hydrophilic polymer such as a functionally activated polyethylene glycol. Also provided are preferred processes for preparing sterile, dry croselinking agents contained within syringes for use in the method of the invention. Methods for sterilization of the croselinking agent include, but are not limited to, sterile filtration, aseptic processing, and e-beam or gamma irradiation Methods for providing augmentation of soft or hard tissue using crosslinked biomaterial compns. prepared according to the method of the invention are also disclosed. A sterile, dry crosslinking agent was prepared by mixing 1500 mg of disfunctionally activated PEG succinimidyl glutarate with 150 mL of water for injection and filtration sterilization using a Durapore filter; 0.5 mL of solution obtained was aliquotted into each of 180 3 cc syringes and lyophilized.

L52 ANSWER 8 OF 8 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1996:195160 CAPLUS

DOCUMENT NUMBER: 124:242408

TITLE: Method of preparing crosslinked biomaterial compositions for use in tissue augmentation

INVENTOR(S): Rhee, Woonza M.; Berg, Richard A.; Rosenblatt, Joel S.; Schroeder, Jacqueline A.; Braga, Larry J.;

Smestad, Thomas L.; Freeman, Abigal PATENT ASSIGNEE(S): Collagen Corporation, USA

SOURCE: Eur. Pat. Appl., 26 pp.

CODEN: EPXXDW
DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 18 PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|----------|----------------------|-------------------|----------|
| EP 697218
EP 697218 | A2
A3 | 19960221
19960529 | EP 1995-112218 | 19950803 |
| R: DE, FR, GB, | IT | | | |
| US 5550187 | A | 19960827 | US 1994-287549 | 19940808 |
| US 5643464 | A | 19970701 | US 1995-497573 | 19950630 |
| PRIORITY APPLN. INFO.: | | | US 1994-287549 A | 19940808 |
| | | | US 1995-497573 A | 19950630 |
| | | | US 1988-274071 B2 | 19881121 |
| | | | US 1989-433441 A2 | 19891114 |
| | | | US 1992-922541 A3 | 19920730 |
| | | | US 1994-198128 A2 | 19940217 |
| | | | US 1994-236769 A2 | 19940502 |

AB The present invention discloses a novel method for preparing crosslinked biomaterial compns. for use in the augmentation of soft or hard tissue. In general, the method comprises mixing a biocompatible polymer, which is preferably collagen, with a sterile, dry crosslinking agent, which is preferably a synthetic hydrophilic polymer such as a functionally activated polyethylene glycol. Also provided are preferred processes for preparing sterile, dry crosslinking agents contained within syringes for use in the method of the invention. Methods for sterilization of the crosslinking agent include, but are not limited to, sterile filtration, aseptic processing, and electron beam or \u03c4-ray irradiation Methods for providing augmentation of soft or hard tissue using crosslinked biomaterial compns. prepared according to the method of the invention are also disclosed. Difunctionally activated PEG succinimidyl glutarate (DSG-PEG) was pelleted with NaCl and the pellet was placed in the barrel of a syringe and mixed with Zyderm I collagen (12 mols of DSG-PEG per mol of collagen) in a syringe and then, a mixture was allowed to crosslink in the syringe. The obtained gel showed a good strength.

=> log h

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L54 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2006:857406 CAPLUS

TITLE: Biomaterials from poly(carboxylic acid) crosslinked

AUTHOR(S): Yang, Yiqi; Reddy, Narendra

CORPORATE SOURCE: Department of Textiles, Clothing and Design and Department of Biological Systems Engineering,

University of Nebraska-Lincoln, Lincoln, NE,

68583-0802, USA SOURCE:

Abstracts of Papers, 232nd ACS National Meeting, San Francisco, CA, United States, Sept. 10-14, 2006 (2006) , CARB-103. American Chemical Society: Washington, D.

CODEN: 69IHRD

DOCUMENT TYPE: Conference; Meeting Abstract; (computer optical disk)

LANGUAGE: English

AB Poly(carboxylic acids) such as citric acid and butanetetracarboxylic acid (BTCA) are inexpensive and non-toxic crosslinking agents that could be used to improve the properties of biomaterials produced from starch. Poly(carboxylic acid) crosslinked starch products are not only relatively inexpensive than starch acetate but have better mech. properties and water stability than similar starch acetate products. Fibers were produced from starch and crosslinked using poly(carboxylic acids) to study the suitability of poly(carboxylic acid) crosslinking as a alternative to starch acetate. Fibers were also produced from starch acetate with various degrees of substitution to compare the properties of crosslinked starch and starch acetate fibers. The fibers produced were tested for their mech. properties and phys. structure. Crosslinked starch fibers had about 300% increase in strength compared to the starch and starch acetate fibers with no change in the elongation of the fibers. Crosslinking also improved the water and biol. resistance of starch fibers. Poly(carboxylic acid) crosslinked starch shows promise to be a cheap alternative to starch acetate for biomaterials.

L54 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2000:529535 CAPLUS

DOCUMENT NUMBER: 133:137057

TITLE: Hot water-resistant gelatin gels useful as

biomaterials INVENTOR(S):

Nagura, Masanobu; Mochizuki, Akira PATENT ASSIGNEE(S): Terumo Corp., Japan

SOURCE: Jpn. Kokai Tokkvo Koho, 4 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
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| | | | | | |
| | JP 2000212286 | A | 20000802 | JP 1999-16038 | 19990125 |
| PRIOF | RITY APPLN. INFO.: | | | JP 1999-16038 | 19990125 |
| ND | The cels are obtain | ed from | delatin sub | stance by crosslinking | with |

polycarboxylic acids under heat and have swelling ratio (Sc) ≤1.0 where $Sc = {Sc(40)/Sc(30)}/{Sh(40)/Sh(30)}$ and 30 and 40 are water temperature in degree (swelling degree Sc and Sh are derived from S = ((Ws-Wd)/Wd; Ws

= weight of gel at equilibrium state; $\ensuremath{\mathtt{Wd}}$ = weight of gel at dry state). Examples of

polycarboxylic acids are succinic acid, citric acid, and adipic acid.

=> log h COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION FULL ESTIMATED COST 15.06 337.42 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL ENTRY SESSION CA SUBSCRIBER PRICE -1.60 -52.80

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 FULL ESTIMATED COST
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 CA SUBSCRIBER PRICE
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4520828 ACID

(CARBOXYLIC OR CARBOXYLICS)

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L55 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER: 2000:529535 CAPLUS
DOCUMENT NUMBER:
                       133:137057
TITLE:
                       Hot water-resistant gelatin gels useful as
                       biomaterials
INVENTOR(S):
                       Nagura, Masanobu; Mochizuki, Akira
PATENT ASSIGNEE(S):
                     Terumo Corp., Japan
SOURCE:
                       Jpn. Kokai Tokkyo Koho, 4 pp.
                       CODEN: JKXXAF
DOCUMENT TYPE:
                       Patent
LANGUAGE:
                       Japanese
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
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     PATENT NO.
    JP 2000212286
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                                                                19990125
PRIORITY APPLN. INFO.:
                                          JP 1999-16038
AB The gels are obtained from gelatin substance by crosslinking with
     polycarboxylic acids under heat and have swelling ratio (Sc)
    \leq 1.0 where Sc = {Sc(40)/Sc(30)}/{Sh(40)/Sh(30)} and 30 and 40 are
     water temperature in degree (swelling degree Sc and Sh are derived from S =
    ((Ws-Wd)/Wd; Ws = weight of gel at equilibrium state; Wd = weight of gel at dry
    state). Examples of polycarboxylic acids are succinic acid, citric
    acid, and adipic acid.
=> s 153 and carboxylic acid
        257842 CARBOXYLIC
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1611629 ACIDS 5028375 ACID

(ACID OR ACIDS) 237609 CARBOXYLIC ACID

(CARBOXYLIC (W) ACID)
L56 6 L53 AND CARBOXYLIC ACID

=> d ibib abs 1-6

L56 ANSWER 1 OF 6 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2008:43481 CAPLUS

TITLE: Biomimetic polymers for tissue engineering
INVENTOR(S): Wang, Yadong; Zern, Blaine; Gumera, Christiane

PATENT ASSIGNEE(S): Georgia Tech Research Corporation, USA

SOURCE: PCT Int. Appl., 49pp.
CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

> PATENT NO. KIND DATE APPLICATION NO. DATE WO 2007-US72946 WO 2008006064 2008006064 A2 20080110 WO 2007-US72946 20070706 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BH, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DO, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, ME, MG, MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RS, RU, SC, SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, LV, MC, MT, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ,

BY, KG, KZ, MD, RU, TJ, TM PRIORITY APPLN. INFO.: US 2006-819219P P 20060707 AB Biodegradable polymers incorporating biomols. and methods of their use are provided. Certain aspects provide biomols, crosslinked with diglycidyl esters. The disclosed compans, have numerous applications including cellular regeneration, wound healing, and cellular differentiation. Thus, a biomimetic polymer PCD was prepared by polymerization of diglycidyl 1,2-cyclohexanedicarboxylate with dopamine in DMF at 90° in a 71% vield. Polymerization of dopamine converted its primary amine to a tertiary amine, which limited the formation of dopaminechrome, the oxidative intermediate to dopamine quinone. This increased the oxidative resistance of the catecholamine, thus minimizing the toxicity associated with dopamine quinone. The ester bond in PCD rendered the polymer biodegradable, with a half-life in phosphate buffered saline solution of approx. 50 days at 37°. Preliminary in vivo biocompatibility studies indicated that PCD did not cause nerve degeneration of fibrous encapsulation when implanted immediately adjacent to rat sciatic nerves. In vitro, neurites

up to 180 µm long began to appear on PCD 3 days after seeding, and grew

L56 ANSWER 2 OF 6 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2006:857406 CAPLUS

TITLE: Biomaterials from poly(carboxylic acid)

crosslinked starch

up to 250 µm after 5 days of culture.

AUTHOR(S): Yang, Yigi; Reddy, Narendra

CORPORATE SOURCE: Department of Textiles, Clothing and Design and

Department of Biological Systems Engineering, University of Nebraska-Lincoln, Lincoln, NE,

68583-0802, USA

Abstracts of Papers, 232nd ACS National Meeting, San SOURCE: Francisco, CA, United States, Sept. 10-14, 2006 (2006)

, CARB-103. American Chemical Society: Washington, D.

CODEN: 69IHRD

DOCUMENT TYPE: Conference; Meeting Abstract; (computer optical disk)

LANGUAGE: English

Poly(carboxylic acids) such as citric acid and butanetetracarboxylic acid (BTCA) are inexpensive and non-toxic crosslinking agents that could be used to improve the properties of biomaterials produced from starch. Poly(carboxylic acid) crosslinked starch products are not only relatively inexpensive than starch acetate but have better mech. properties and water stability than similar starch acetate products. Fibers were produced from starch and crosslinked using poly(carboxylic acids) to study the suitability of poly(carboxylic acid) crosslinking as a alternative to starch acetate. Fibers were also produced from starch acetate with various degrees of substitution to compare the properties of crosslinked starch and starch acetate fibers. The fibers produced were tested for their mech. properties and phys. structure. Crosslinked starch fibers had about 300% increase in strength compared to the starch and starch acetate fibers with no change in the elongation of the fibers. Crosslinking also improved the water and biol. resistance of starch fibers. Poly(carboxylic acid) crosslinked starch shows promise to be a cheap alternative to starch acetate for

L56 ANSWER 3 OF 6 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2004:183216 CAPLUS DOCUMENT NUMBER: 140:223219

biomaterials.

TITLE:

Controllable Surface Modification of Poly(lactic-co-glycolic acid) (PLGA) by Hydrolysis or

Aminolysis I: Physical, Chemical, and Theoretical

AUTHOR(S): Croll, Tristan I.; O'Connor, Andrea J.; Stevens, Geoffrey W.; Cooper-White, Justin J.

Department of Chemical and Biomolecular Engineering, CORPORATE SOURCE:

The University of Melbourne, Melbourne, 3010,

Australia

SOURCE: Biomacromolecules (2004), 5(2), 463-473

CODEN: BOMAF6; ISSN: 1525-7797 American Chemical Society

DOCUMENT TYPE: Journal

PUBLISHER:

LANGUAGE: English

AB While biodegradable, biocompatible polyesters such as poly

(lactic-co-glycolic acid) (PLGA) are popular materials for the manufacture of tissue engineering scaffolds, their surface properties are not particularly suitable for directed tissue growth. Although a number of approaches to chemical modify the PLGA surface have been reported, their

applicability to soft tissue scaffolds, which combine large vols., complex shapes, and extremely fine structures, is questionable. In this paper, we describe two wet-chemical methods, base hydrolysis and aminolysis, to introduce useful levels of carboxylic acid or primary and secondary amine groups, resp., onto the surface of PLGA with minimal degradation The

effects of temperature, concentration, pH, and solvent type on the kinetics of these

reactions are studied by following changes in the wettability of the PLGA

using contact angle measurements. In addition, the treated surfaces are studied using XPS to determine the effect on the surface chemical structure. Furthermore, we show using XPS anal. that these carboxyl and amine groups are readily activated to allow the covalent attachment of biol. macromols. REFERENCE COUNT: 55 THERE ARE 55 CITED REFERENCE AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L56 ANSWER 4 OF 6 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2000:529535 CAPLUS

DOCUMENT NUMBER: 133:137057

TITLE: Hot water-resistant gelatin gels useful as

biomaterials
INVENTOR(S): Nagura, Masanobu; Mochizuki, Akira

PATENT ASSIGNEE(S): Terumo Corp., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 4 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| | PATENT | NO. | KIND | DATE | API | PLICATION | NO. | DATE |
|-------|---------|-------------|------|----------|-----|-----------|-----|----------|
| | | | | | | | | |
| | | 0212286 | A | 20000802 | | 1999-1603 | | 19990125 |
| PKIOK | ITT API | PLN. INFO.: | | | JP | 1999-1603 | 38 | 19990125 |

AB The gels are obtained from gelatin substance by crosslinking with polycarboxylic acids under heat and have swelling ratio (Sc) ≤1.0

where Sc = $\{Sc(40)/Sc(30)\}/\{Sh(40)/Sh(30)\}$ and 30 and 40 are water temperature in degree (swelling degree Sc and Sh are derived from S = $\{(Ws-Wd)/Wd; Ws = weight of gel at equilibrium state; Wd = weight of gel at dry state).$

Examples of

polycarboxylic acids are succinic acid, citric acid, and adipic acid.

L56 ANSWER 5 OF 6 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1996:660894 CAPLUS

DOCUMENT NUMBER: 125:285011

TITLE: Use of hydrophobic crosslinking agents to prepare

crosslinked biomaterial implants

INVENTOR(S): Rhee, Woonza M.

PATENT ASSIGNEE(S): Collagen Corporation, USA SOURCE: Eur. Pat. Appl., 26 pp.

Eur. Pat. Appl., 26 pp. CODEN: EPXXDW

Patent

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|--------|-----------|------------------|------------|
| | | | | |
| EP 732109 | A1 | 19960918 | EP 1996-102366 | 19960216 |
| R: AT, CH, DE, | FR, GB | , IT, LI, | NL, SE | |
| CA 2165728 | A1 | 19960915 | CA 1995-2165728 | 19951220 |
| JP 09249751 | A | 19970922 | JP 1996-58138 | 19960314 |
| US 6962979 | B1 | 20051108 | US 1999-344230 | 19990625 |
| US 2004121951 | A1 | 20040624 | US 2003-448246 | 20030528 |
| US 7129209 | B2 | 20061031 | | |
| US 2005154125 | A1 | 20050714 | US 2004-997246 | 20041123 |
| JP 2006181389 | A | 20060713 | JP 2006-66762 | 20060310 |
| PRIORITY APPLN. INFO.: | | | US 1995-403358 A | 19950314 |
| | | | JP 1996-58138 A | 3 19960314 |

US 1997-987467 B1 19971209 US 1999-344230 A1 19990625

AB Novel crosslinked biomaterial compns. are prepared using hydrophobic polymers as a crosslinking agent. Preferred hydrophobic polymers are those that contain two or more reactive succinimidyl groups, including disuccinimidyl suberate (I), bis(sulfosuccinimidyl) suberate, and dithiobis(succinimidyl propionate). Crosslinked biomaterial compns. prepared using mixts, of hydrophobic and hydrophilic crosslinking agents are also disclosed. The compns. of the present invention can be used to prepare formed implants for use in a variety of medical applications. Thus, 1.0 mL of 35 mg/mL collagen was mixed with 3 mg I in a syringe and incubated at 37° for 16 h. The crosslinked collagen material was extruded out of the plunger end of the syringe and the resulting crosslinked cylindrical gels were then sectioned into 5 mm thick disks. The solubilization of crosslinked collagen in trypsin solution and oxidative degradation in 3% H202 was 7 and 14 days, resp. After implantation of the crosslinked collagen in rats for 90 days it had a discrete, football-shaped, bolus-like configuration, whereas noncrosslinked formulation was present as a more diffuse mass.

L56 ANSWER 6 OF 6 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1996:254782 CAPLUS

DOCUMENT NUMBER: 124:325312

TITLE: Crosslinking of dermal sheep collagen using a

water-soluble carbodiimide

Olde Damink, L. H. H.; Dijkstra, P. J.; van Luyn, M. AUTHOR(S): J. A.; van Wachem, P. B.; Nieuwnehuis, P.; Feijen, J. CORPORATE SOURCE: Dep. Chem. Technol., Univ. Twente, Enchede, 7500 AE,

Neth.

Biomaterials (1996), 17(8), 765-73 SOURCE: CODEN: BIMADU; ISSN: 0142-9612

PUBLISHER: Elsevier DOCUMENT TYPE: Journal

LANGUAGE: English AB A crosslinking method for collagen-based biomaterials was developed

using the water-soluble carbodiimide 1-ethyl-3-(3-dimethyl aminopropyl)carbodiimide hydrochloride (EDC). Crosslinking using EDC involves the activation of carboxylic acid groups to give O-acvlisourea groups, which form crosslinks after reaction with free amine groups. Treatment of dermal sheep collagen (DSC) with EDC (E-DSC) resulted in materials with an increased shrinkage temperature (Ts) and a decreased free amine group content, showing that crosslinking occurred. Addition of N-hydroxysuccinimide to the EDC-containing crosslinking solution (E/N-DSC) increased the rate of crosslinking. Crosslinking increased the Ts of non-crosslinked DSC samples from 56 to 73 °C for E-DSC and to 86°C for E/N-DSC samples, resp. For both crosslinking methods a linear relation between the decrease in free amine group content and the increase in Ts was observed The tensile strength and the high strain modulus of E/N-DSC samples decreased upon crosslinking from 18 to 15 MPa and from 26 to 16 MPa, resp. The elongation at break of E/N-DSC increased upon crosslinking from 142 to 180%.

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=> s 146 and citric acid
         94137 CITRIC
             2 CITRICS
         94139 CITRIC
                 (CITRIC OR CITRICS)
       4520828 ACID
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1611629 ACIDS

L29

0 S E3/RACT

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L57
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                E CITRIC ACID/CN
                E MALIC ACID/CN
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                E CITRIC ACID/CN
L2
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                E OXALACETIC ACID/CN
              1 S E3
                E CITRIC ACID/CN
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                E ACONITIC ACID/CN
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               E MALATE
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L9
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L12
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L13
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L33
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L35
           800 S L6/RACT
L36
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L37
          184 S L15/RACT
L38
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1.39
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L42
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L43
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1.45
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L46
         72274 S CROSSLINKING AGENT
L47
        595653 S SULFATE
L48
          2066 S L46 AND L47
L49
              8 S CHRONDROITIN SULFATE
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L50
         13474 S (CHONDROITIN SULFATE OR "CHONDROITIN, HYDROGEN SULFATE")
L51
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L52
             8 S L51 AND BIOMATERIAL
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L54
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L55
             1 S L53 AND POLYCARBOXYLIC ACID
L56
             6 S L53 AND CARBOXYLIC ACID
           617 S L46 AND CITRIC ACID
L57
=> s 157 and collagen
         95914 COLLAGEN
        68797 COLLAGENS
        107638 COLLAGEN
                 (COLLAGEN OR COLLAGENS)
L58
           17 L57 AND COLLAGEN
=> d ibib abs 1-17
L58 ANSWER 1 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                        2007:1454937 CAPLUS
DOCUMENT NUMBER:
                        148:85879
TITLE:
                        Collagen cross-linking agents such as bioflavonoid
                        compounds, grape seed extract, casein
                        phosphopeptide-amorphous calcium phosphate, or iridoid
                        compounds, on dental restorative treatment and
```

preventive dentistry INVENTOR(S): Bedran-Russo, Ana K.

PATENT ASSIGNEE(S): The Board of Trustees of the University of Illinois,

SOURCE: PCT Int. Appl., 45pp. CODEN: PIXXD2

Patent DOCUMENT TYPE: Enalish

LANGUAGE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| PA: | TENT | NO. | | | KIN | D | DATE | | | APPL | ICAT | ION | NO. | | D. | ATE | |
|-----|------|------|-----|-----|-------------|-----|------|-----|-----|------|------|------|-----|-----|----------|-----|-----|
| | | | | | | - | | | | | | | | | - | | |
| WO | 2007 | 1468 | 41 | | A2 20071221 | | | | | WO 2 | 007- | US70 | 809 | | 20070608 | | |
| | W: | ΑE, | AG, | AL, | AM, | AT, | AU, | AZ, | BA, | BB, | BG, | BH, | BR, | BW, | BY, | BZ, | CA, |
| | | CH, | CN, | CO, | CR, | CU, | CZ, | DE, | DK, | DM, | DO, | DZ, | EC, | EE, | EG, | ES, | FI, |
| | | GB, | GD, | GE, | GH, | GM, | GT, | HN, | HR, | HU, | ID, | IL, | IN, | IS, | JP, | KE, | KG, |
| | | KM, | KN, | KP, | KR, | KZ, | LA, | LC, | LK, | LR, | LS, | LT, | LU, | LY, | MA, | MD, | ME, |
| | | MG, | MK, | MN, | MW, | MX, | MY, | MZ, | NA, | NG, | NI, | NO, | NZ, | OM, | PG, | PH, | PL, |
| | | PT, | RO, | RS, | RU, | SC, | SD, | SE, | SG, | SK, | SL, | SM, | SV, | SY, | TJ, | TM, | TN, |
| | | | | | | | | | | | | | | | | | |

TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, LV, MC, MT, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

PRIORITY APPLN. INFO .: US 2006-812664P P 20060609 US 2007-918640P P 20070316

ΔR The invention relates to the development of compns. and methods for increasing the amount of collagen crosslinking in a mammalian tissue. A typical composition as described herein includes at least one crosslinking agent such as a bioflavonoid compound (e.g., proanthocyanidin), a grape seed extract, a casein phosphopeptide-amorphous calcium phosphate, or an iridoid compound (e.g., genipin) in an amount effective for increasing collagen crosslinking in the mammalian tissue in a pharmaceutically acceptable carrier. A typical method for increasing the amount of collagen crosslinking in dentin in a mammalian tooth includes the steps of preparing the surface of the tooth to be treated; and applying a composition including at least one of a bioflavonoid compound, a grape seed extract, a casein phosphopeptide-amorphous calcium phosphate, and an iridoid compound in a pharmaceutically acceptable carrier to the tooth surface for a time period of 0.0001 h to about 4 h. In some embodiments, two or more crosslinking agents are included in the compns. described herein. The compns. and methods as described herein are particularly useful for applying to dentin in a mammalian tooth requiring a restorative procedure for improving the mech. properties of restoration interfaces to withstand degradation over time. Compns. containing one of the collagen crosslinking agents as described herein were applied to dentin collagen and resulted in a significant improvement in ultimate tensile strength indicating the value of these compns. in restorative dentistry. The compns. and methods described herein will also find use in preventive dentistry applications, and can be applied to sound dentin, caries-affected dentin, and dentin that is impaired, weak, or degraded in any way. Thus, the effect of three biocompatible collagen crosslinking agents on the ultimate tensile strength (UTS) of dentin was tested: 5% glutaraldehyde (GD); 0.5% proanthocyanidin PBS solution (PA); and 0.625% genipin PBS solution (GE). A highly significant increase in UTS values was observed after PA dentin treatment, compared to the control and the other two crosslinking agents: the increase of almost 70% and 110% in the UTS values after PA treatment during 4 and 40 h, resp., indicates a

great potential of the agent to induce crosslinks in the dentin collagen.

L58 ANSWER 2 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2007:813931 CAPLUS

DOCUMENT NUMBER: 147:183443

TITLE: Protein compositions containing water soluble salts, and their articles with improved mechanical properties

INVENTOR(S): Hirase, Ryuji; Nakagawa, Kazuharu; Kubo, Junichi

PATENT ASSIGNEE(S): Hyogo Prefecture, Japan; Ako Kasei Co., Ltd.

SOURCE: Jpn. Kokai Tokkyo Koho, 12pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE:

Japanese FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE | | | | |
|------------------------|--------|-------------|-------------------------|----------|--|--|--|--|
| | | | | | | | | |
| JP 2007186556 | A | 20070726 | JP 2006-4289 | 20060112 | | | | |
| PRIORITY APPLN. INFO.: | | | JP 2006-4289 | 20060112 | | | | |
| AB The compns., which | can be | formed into | shaped articles such as | films, | | | | |

sheets, yarns, and rods for industrial materials or foods, contain proteins and water-soluble inorg. salt hydrates or water-soluble organic acid

metal

salt hydrates. The compns. may also contain crosslinking agents.

Thus, an aqueous solution containing 5 g JS-110 (gelatin) was mixed with 1.0 g MgCl2.6H2O, cast in a mold, and dried at 50° for .apprx.48 h to

give a film showing maximum stress 8.3 MPa and elongation at break 256.8%.

L58 ANSWER 3 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2007:729205 CAPLUS

DOCUMENT NUMBER: 147:156885 TITLE:

Production of chiral materials using crystallization inhibitors

INVENTOR(S):

Valluzzi, Regina; Liu, Liya

Evolved Nanomaterial Sciences, Inc., USA PATENT ASSIGNEE(S): SOURCE: PCT Int. Appl., 53pp.

CODEN: PIXXD2 Patent

DOCUMENT TYPE: LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2 PATENT INFORMATION:

| PA' | TENT | | KIND DATE | | | | | APPL | ICAT | | DATE | | | | | | | |
|-----|---------------|--|--|--|--|--|--|--|---------------------------------|---------------------------------|---------------------------------|--------------------------|--------------------------|--------------------------|--------------------------|--------------------------|--------------------------|--|
| | 2007 | | 09 | | A2
A3 | | 2007
2007 | 0705
0913 | | WO 2 | 006- | US 48 | 312 | | 20061219 | | | |
| | W: | AE,
CN,
GE,
KP,
MN,
RS, | AG,
CO,
GH,
KR,
MW,
RU, | AL,
CR,
GM,
KZ,
MX,
SC, | AM,
CU,
GT,
LA,
MY,
SD, | AT,
CZ,
HN,
LC,
MZ,
SE, | AU,
DE,
HR,
LK,
NA,
SG, | AZ,
DK,
HU,
LR,
NG,
SK, | DM,
ID,
LS,
NI,
SL, | DZ,
IL,
LT,
NO,
SM, | EC,
IN,
LU,
NZ,
SV, | EE,
IS,
LV,
OM, | EG,
JP,
LY,
PG, | ES,
KE,
MA,
PH, | FI,
KG,
MD,
PL, | GB,
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LU,
CM,
MW,
RU, | CY,
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MZ,
TJ, | CZ,
MC,
GN,
NA,
TM, | NL,
GQ,
SD,
AP, | DK,
PL,
GW,
SL,
EA, | EE,
PT,
ML,
SZ,
EP, | ES,
RO,
MR,
TZ,
OA | SE,
NE,
UG, | SI,
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AM, | BF,
BW,
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GH,
BY, | |
| US | JS 2007255042 | | | A1 | | 20071101 | | | US 2006-641344 | | | | | 20061219 | | | | |

PRIORITY APPLN. INFO.:

US 2005-751545P P 20051219 US 2006-785669P P 20060324

A method is disclosed for producing a chiral gel. A polymer including chiral monomers, such as a protein, is dissolved to generate a sol, which is optionally dialyzed. The sol is contacted with a crystallization inhibitor that allows it to form a gel. The gel in wet or dried form is useful for performing chiral sepns.

L58 ANSWER 4 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2006:1349735 CAPLUS

DOCUMENT NUMBER: 146:87679

TITLE: Solid-liquid mixing two-component-type biodegradable

medical adhesive materials

INVENTOR(S): Taguchi, Satoshi; Kakinoki, Sachiro; Tanaka, Junzo;

Saito, Hiroshi

PATENT ASSIGNEE(S): National Institute of Materials Science, Japan;

Furuuchi Kagaku Co., Ltd. SOURCE: Jpn. Kokai Tokkyo Koho, 11pp.

CODEN: JKXXAF DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | KIND | DATE . | APE | LICATION NO. | D | ATE |
|------------------------|---------|--------------|-----|---------------|---------|---------|
| | | | | | | |
| JP 2006346049 | A | 20061228 | JΡ | 2005-174414 | 2 | 0050614 |
| PRIORITY APPLN. INFO.: | | | JΡ | 2005-174414 | 2 | 0050614 |
| AB The invention relat | es to a | two-agent-tv | pe | biodegradable | medical | adhesiv |

material consisting of a liquid adhesive agent and a powder hardening agent for use by mixing together at the usage. The liquid adhesive agent contains water, biodegradable polymer, and a solution with metal ions which interacts with the biodegradable polymer through electrostatic effect or cheating effect, or the liquid adhesive agent contains a biodegradable polymer dissolved in a buffer solution The powder agent contains a di- or tri-carboxylic acid derivative whose at least 2 carboxylic groups are modified by electron-attracting groups, e.g. succinimidyl, sulfosuccinimidyl, maleimidyl, phthalimidyl, imidazolyl, nitrophenyl, and tresyl. The biodegradable polymer and the hardening agent are reacted by mixing to form a crosslinked adhesive material. For example, human-derived albumin in phosphate buffer solution was mixed with citric acid

N-hydroxysuccinimide derivative to form an adhesive material.

L58 ANSWER 5 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2004:428813 CAPLUS

DOCUMENT NUMBER: 140:412344

TITLE: Pharmaceutical compositions and dosage forms for buccal and sublingual delivery of tizanidine and

methods of administering tizanidine sublingually or

bucally

Lerner, Itzhak E.; Flashner-Barak, Moshe; Rosenberger, INVENTOR(S):

Vered

PATENT ASSIGNEE(S): Teva Pharmaceutical Industries Ltd., Israel: Teva

Pharmaceutical USA, Inc. SOURCE: PCT Int. Appl., 27 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

```
PATENT NO.
                     KIND DATE APPLICATION NO. DATE
     WO 2004043431 A1 20040527 WO 2003-US35002 20031103
          W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
               CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
                GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
                LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM,
                PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN,
                TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
          RW: BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ,
                BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE,
                ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK,
                TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
     CA 2505861 A1 20040527 CA 2003-2505861 20031103
AU 2003287488 A1 20040603 AU 2003-287488 20031103
     AU 2003287488 AI 20040603 AU 2003-287488 20031103
AU 2003287488 B2 20070405
US 2004122065 AI 20040624 US 2003-699991 20031103
BR 2003015482 A 20050823 BR 2003-15482 20031103
EP 1567124 AI 20050831 EP 2003-781729 20031103
          R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
               IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
     CN 1738600 A 20060222 CN 2003-80108649 20031103

JP 2006508122 T 20060309 JP 2004-551688 20031103

MX 2005PA05038 A 20050701 MX 2005-PA5038 20050511

MITY APPLN. INFO.: US 2002-425326P P 20021112

WO 2003-US355002 W 20031US35502
PRIORITY APPLN. INFO.:
     Sublingual and buccal administration of the muscle spasm suppressor
```

tizanidine increase its bioavailability by avoiding first-pass metabolism in the liver and reduce the inter-patient variation in bioavailability. REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L58 ANSWER 6 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2004:354279 CAPLUS

DOCUMENT NUMBER: 140:344534

TITLE: Antiinflammatory sheet packs containing glycyrrhetinic

acid, glycyrrhizic acid, or their esters

INVENTOR(S): Hinobu, Kimiko; Iida, Norio PATENT ASSIGNEE(S): Lion Corp., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 38 pp. INVENTOR(S):

CODEN: JKXXAF

DOCUMENT TYPE: Patent

LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|------|----------|-----------------|----------|
| | | | | |
| JP 2004131383 | A | 20040430 | JP 2002-184592 | 20020625 |
| PRIORITY APPLN. INFO.: | | | JP 2002-184592 | 20020625 |
| | | | | |

A sheet pack comprises a support and an aqueous pressure-sensitive adhesive containing crosslinked polyacrylate matrix, ≥1 inflammation inhibitors selected from glycyrrhetinic acid, glycyrrhizic acid, and their esters, and H2O. The pack is less skin-irritating and conditions skin damaged by drying, allergy, UV, sunburn, etc. A laminate of a polyethylene film and a thermally-bonded polyester nonwoven fabric was coated with an adhesive composition containing poly(acrylic acid) (mol. weight 100,000-300,000) 3, poly(acrylic acid) (mol. weight 500,000-1,200,000) 2, Na polyacrylate 1.5, CM-cellulose Na 3, glycerin 15, 70% sorbitol solution 10, polyoxyethylene

lauryl ether 1, methylparaben 0.2, glycyrrhetinic acid 0.1, Aloe extract 0.1, alginic acid 0.5, bentonite 2, synthetic hydrotalcite 0.1, Al glycinate 0.1, Na edetate 0.01%, and H2O balance and covered with a PET film to give a sheet pack. The pack showed good face-moisturizing effect in female volunteers.

L58 ANSWER 7 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2004:252481 CAPLUS

DOCUMENT NUMBER: 140:287718

TITLE: Preparation of biological low-molecular weight carboxylic acid derivatives as crosslinking agents

for biopolymers
INVENTOR(S): Taguchi, Tetsushi; Kobayashi, Naotoshi; Tanaka, Junzo;

Saito, Hiroshi

PATENT ASSIGNEE(S): National Institute for Materials Science, Japan; Furuuchi Chemical Corporation

SOURCE: PCT Int. Appl., 20 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: Japanese FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| | PATENT NO. | | | | | KIND DATE | | | APPLICATION NO. | | | | | DATE | | | | |
|-------|------------|-------|-------|-------|-------|-----------|-----|------|-----------------|-------|------|------|-------|------|-----|------|-------|-----|
| | WO | 2004 | 0246 | 36 | | A1 | | 2004 | 0325 | WO | 200 | 03-3 | JP116 | 669 | | 20 | 0030 | 911 |
| | | W: | CA, | CN, | US | | | | | | | | | | | | | |
| | | RW: | AT, | BE, | BG, | CH, | CY, | CZ, | DE, | DK, E | Ε, Ι | ES, | FI, | FR, | GB, | GR, | HU, | IE, |
| | | | IT, | LU, | MC, | NL, | PT, | RO, | SE, | SI, S | к, : | TR | | | | | | |
| | JP | 2004 | 09956 | 52 | | A | | 2004 | 0402 | JP | 200 | 02-2 | 26598 | 82 | | 20 | 0020 | 911 |
| | CA | 2499 | 606 | | | A1 | | 2004 | 0325 | CA | 200 | 03-2 | 24996 | 606 | | 20 | 0030 | 911 |
| | EP | 1548 | 004 | | | A1 | | 2005 | 0629 | EP | 200 | 03- | 7954: | 11 | | 20 | 0030 | 911 |
| | | R: | AT, | BE, | CH, | DE, | DK, | ES, | FR, | GB, G | R, : | IT, | LI, | LU, | NL, | SE, | MC, | PT, |
| | | | ΙE, | SI, | FI, | RO, | CY, | TR, | BG, | CZ, E | Ε, Ι | HU, | SK | | | | | |
| | CN | 1681 | 780 | | | A | | 2005 | 1012 | CN | 200 | 03-8 | 3215 | 40 | | 20 | 0030 | 911 |
| | US | 2006 | 1289 | 18 | | A1 | | 2006 | 0615 | US | 200 | 05-5 | 2769 | 94 | | 20 | 0051 | 101 |
| PRIOR | RITY | Y APP | LN. : | INFO | . : | | | | | JP | 200 | 02-2 | 26591 | 82 | I | 1 20 | 0020 | 911 |
| | | | | | | | | | | WO | 200 | 03-3 | JP116 | 669 | V | 7 20 | 0030 | 911 |
| AB | Ιt | is p | ointe | ed or | at th | hat 1 | he | exis | ting | cross | lin | king | g age | ents | and | cond | dens. | ing |

agents having been employed in biol. adhesives and in treating medical devices such as cardiac valves, which are non-natural products synthesized artificially, are not metabolized in vivo and exhibit toxicity to living bodies. Therefore, these products can be used only in a restricted amount and for limited purposes in the clin. field. It is intended to provide biol. low-mol. weight derivs. obtained by modifying a carboxyl group of a biol. low-mol. weight compound such as malic acid, oxalacetic acid, citric acid, cis-aconitic acid, and 2-ketoglutaric acid with N-hydroxysuccinimide, N-hydroxysulfosuccinimide or derivs. thereof and crosslinked high-mol. weight compds. obtained by crosslinking various high-mol. weight compds. such as polysaccharides and proteins with the use of the above derivative Gels containing biopolymers and crosslinking acents are

crosslinked directly at disease sites and applied as bioadhesives, hemostatics, vascular embolus agents, encapsulants for aneurysm. Crosslinked biopolymers are used as adhesion inhibitors, base materials for tissue reqeneration, and drug carriers.

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L58 ANSWER 8 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2004:183096 CAPLUS

DOCUMENT NUMBER: 140:234396

TITLE: Antibodies and other binding agents specific to

thrombospondin fragments for diagnosis of cancer and other diseases

INVENTOR(S): Williams, Kevin J.

PATENT ASSIGNEE(S): USA

SOURCE: PCT Int. Appl., 76 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent English

LANGUAGE: FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

| | | | | | | APPLICATION NO. | | | | | | | | | | | |
|------|---|------|-----|-----|-----|-----------------|------|------------------|-----------------|------|------|------|-----|------------|-----|------|-----|
| WO : | | 0189 | 95 | | | | | WO 2003-US326023 | | | | | | | | | |
| | W: | ΑE, | AG, | AL, | AM, | ΑT, | AU, | ΑZ, | | | | | | | | | |
| | | GM, | HR, | HU, | ID, | IL, | IN, | DM,
IS, | JP, | KE, | KG, | KP, | KR, | KZ, | LC, | LK, | LR, |
| | | | | | | | | MG,
SC, | | | | | | | | | |
| | DIT. | TR, | TT, | TZ, | UA, | UG, | US, | UZ,
SD, | VC, | VN, | YU, | ZA, | ZM, | zw | | | |
| | KW: | KG, | ΚZ, | MD, | RU, | ΤJ, | TM, | ΑT, | BE, | BG, | CH, | CY, | CZ, | DE, | DK, | EE, | ES, |
| | | | | | | | | IT,
GA, | | | | | | | | | |
| | | | | | | | | 0318 | | | | | | | | | |
| | | | | | | | | 0304
0311 | | | | | | | | | |
| | 1572 | 225 | | | A2 | | 2005 | 0914 | | EP 2 | 003- | 7931 | 49 | | 2 | 0030 | 820 |
| | R: | | | | | | | FR,
MK, | | | | | | | | | PT, |
| | | | | | | | | 0324 | | | | | | | | | |
| | US 2006257947
PRIORITY APPLN. INFO.: | | | | | | 2006 | 1110 | | US 2 | | 4054 | 94P | P 20020823 | | | |
| | | | | | | | | | WO 2003-US26023 | | | | | | | | |

The invention relates to thrombospondin fragments found in plasma, their use or use of portions thereof in diagnostic methods, as method calibrators, method indicators, and as immunogens, and as analytes for methods with substantial clin. utility; and their detection in plasma or other bodily fluids for purpose of diagnostic methods, especially for cancer. The thrombospondin fragments include fibronectin-binding domain, procollagen homol. region, type 1 and 2 repeats, amino-terminal domain, and heparin-binding domain. The antibodies are useful for diagnosis of cancer, metastasis, renal failure, atopic dermatitis, acute vasculitis, asthma, diabetes mellitus, rheumatoid arthritis, myocaridal infarction, inlfammatory disease, blood clotting conditions, etc.

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L58 ANSWER 9 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                       2004:142611 CAPLUS
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DOCUMENT NUMBER: 140:187393

TITLE: Composite matrix containing chitosan derivatives for microcapsules

INVENTOR(S): Chen, Yuan Han; Yeh, Ming Hsi; Lai, Huey Min PATENT ASSIGNEE(S): Industrial Technology Research Institute, Taiwan; Chiu, Kuo-Cheng

U.S. Pat. Appl. Publ., 8 pp. SOURCE:

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

weight% to

PATENT NO. KIND DATE APPLICATION NO. DATE US 2004033265 A1 20040219 US 2002-329712 20021227 RITY APPEN. INFO: TW 2001-90133359 A 20011231 PRIORITY APPLN. INFO.:

AB A method for preparation of a composite matrix containing chitosan derivs., comprising the steps of: (i) providing an anionic chitosan derivative solution (A); (ii) providing a cationic polysaccharide solution (B); and (iii) mixing solution (A) and solution (B) to form microcapsules. Metallic ion crosslinking agent and/or natural protein solution can be added optionally to adjust the mech. strength of the shell and the interior physic state of the microcapsules. For example, 2 weight% N,O-carboxymethyl chitosan (NOCC) solution was dropped into the stirring mixture containing 1

4 weight% of chitosan dissolved in 1 weight% acetic acid solution, 1 weight% collagen dissolved in 1 weight% acetic acid solution, and 1M to 5M of calcium chloride solution, wherein the weight ratio of chitosan to collagen to calcium ion is 6:1:3, 9:2:9 or 3:1:6. The NOCC converses to microcapsules immediately when it contacts the mixture The diameter of the microcapsule could be adjusted by controlling size of the droplet. The diameter of thus obtained microcapsules ranges from 8 mm to 0.2 mm. The shell of the microcapsules increases with soaking time, and microcapsules with liquid interior will form ultimately.

L58 ANSWER 10 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2004:59579 CAPLUS

DOCUMENT NUMBER: 140:99664

TITLE: Preparation of a biodegradable thermal-sensitive gel

system

INVENTOR(S): Chem, Yuan-han; Yeh, Ming-hsi; Lai, Huey-min
PATENT ASSIGNEE(S): Industrial Technology Research Institute, Taiwan
SOURCE: U.S. Pat. Appl. Publ., 11 pp.

CODEN: USXXCO DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. PATENT NO. KIND DATE APPLICATION NO. DATE US 2004013733 A1 20040122 US 2002-330085 20021230 TW 245634 B 20051221 TW 2002-91124213 20021021 PRIORITY APPLN. INFO.: TW 2001-90132965 A 20011228 TW 2002-91124213 A 20021021

The present invention relates to a biodegradable thermal-sensitive gel system, which comprises at least one polysaccharide solution, at least one electrolytic salt, and at least one buffer solution for adjusting pH. A natural protein as well as a crosslinking agent can be added to the gel system optionally. Said gel system is liquid at room temperature (25°) and solidifies at or above 37°. The present invention also relates to a process for preparing said gel system, and a use for drug releasing carrier. For example, a gel system with natural proteins was prepared by adding 4 mL of 4 weight% chitosan (in 1 weight% acetic acid) and 1 mL of 1 weight%

collagen (in 1 weight% acetic acid) to 3 mL of PBS (pH 7.6) at room temperature with stirring, followed by 1 mL of 56 weight% glycerol-phosphate and 1 mL of

 $0.5~\rm M\, AHCO3$ to adjust the pH value of the solution to 7.2. The product thus obtained is liquid and will solidify while the temperature rises to $37\,^{\rm c},$ which needs about 3 min.

L58 ANSWER 11 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2002:122835 CAPLUS

DOCUMENT NUMBER: 136:172843

TITLE: Method for the production of chitosan-based films with enhanced cell adhering capacity, resulting product and

applications

INVENTOR(S): Lopez Lacomba, Jose Luis; Garcia Cantalejo, Jesus Manuel; Sanz Casado, Jose Vicente; Ramos, Viviana

Monica

PATENT ASSIGNEE(S): Osfarma, S.L., Spain

SOURCE: PCT Int. Appl., 69 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent

LANGUAGE: Spanish FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| | | TENT | | | | | | DATE | | | | ICAT | | | | | ATE | | |
|-----|-------|--------------|------|------|-----|-----|-----|------|------|-----|------|-------|------|-----|------|-----|--------|-------|--|
| | | 2002 | | | | | | | | | | | | | | | | | |
| | WO | 2002 | 0117 | 82 | | A8 | | 2002 | 0711 | | | | | | | | | | |
| | | W: | AE, | AG, | AL, | AM, | AT, | AU, | AZ, | BA, | BB, | BG, | BR, | BY, | BZ, | CA, | CH, | CN, | |
| | | | | | | | | | | | | EE, | | | | | | | |
| | | | | | | | | | | | | KG, | | | | | | | |
| | | | | | | | | | | | | MW, | | | | | | | |
| | | | | | | | | | | | | TM, | | | | | | | |
| | | | | | | ZA. | | | | | | | | | | | | , | |
| | | RW: | GH. | GM. | KE. | LS. | MW. | MZ. | SD. | SL. | SZ. | TZ, | UG. | ZW. | AT. | BE. | CH. | CY. | |
| | | | | | | | | | | | | LU, | | | | | | | |
| | | | | | | | | | | | | ML, | | | | | | , | |
| | ES | 2169 | | | | | | | | | | 2000- | | | | | | 810 | |
| | ES | 2169 | | | | | | 2003 | | | | | | | | | | | |
| | AU | 2001 | 0821 | 53 | | A5 | | 2002 | 0218 | | AU 2 | 2001- | 8215 | 3 | | 2 | 20010 | 810 | |
| | EP | 1308 | 177 | | | A1 | | 2003 | 0507 | | EP 2 | 2001- | 9607 | 53 | | - 2 | 20010 | 810 | |
| | | 1308 | | | | | | | | | | | | | | | | | |
| | | | | | | | | | | | GR. | IT, | LI. | LU. | NL. | SE. | MC. | PT. | |
| | | | | | | | | RO, | | | | | | | | | | | |
| | BR | 2001 | 0131 | 06 | | A | | 2003 | 0715 | | BR 2 | 2001- | 1310 | 6 | | 2 | 20010 | 810 | |
| | JP | 2004
2951 | 5056 | 78 | | T | | 2004 | 0226 | | JP 2 | 2002- | 5171 | 14 | | 2 | 20010 | 810 | |
| | AT | 2951 | 90 | | | T | | 2005 | 0515 | | AT 2 | 2001- | 9607 | 53 | | 2 | 20010 | 810 | |
| | ES | 2246 | 337 | | | Т3 | | 2006 | 0216 | | ES 2 | 2001- | 1960 | 753 | | 2 | 20010 | 810 | |
| | US | 2003 | 1241 | 72 | | A1 | | 2003 | 0703 | | US 2 | 2003- | 3648 | 27 | | 2 | 0030 | 210 | |
| PR | IORIT | Y APP | LN. | INFO | . : | | | | | | ES 2 | -000 | 2057 | | | A 2 | 0000 | 810 | |
| | | | | | | | | | | | WO 2 | 2001- | ES32 | 2 | 1 | W 2 | 20010 | 810 | |
| 3 D | 0.0 | | | | | 1 | | | c | | | | | | £11. | | Lake 4 | 14-4- | |

AB Said method generally involves forming a chitosan-based film; activating the cell adhering capacity by drying the stabilized film and washing. The film can also be activated biol. by fixing a substance with biol. activity. The resulting films exhibit enhanced cell adhering capacity and are optionally biol. activated. Said films can be used to induce biol. activity in a receiver organism and/or enhance osteo-integration of implants used in odontol. or traumatol. and/or

regenerate bone tissue.
REFERENCE COUNT: 9

THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

2000:508939 CAPLUS ACCESSION NUMBER: 133.94623

DOCUMENT NUMBER: Manufacture of medical collagen sponge TITLE:

INVENTOR(S): Zhan, Lifen

PATENT ASSIGNEE(S): Peop. Rep. China

SOURCE: Faming Zhuanli Shenging Gongkai Shuomingshu, 6 pp.

CODEN: CNXXEV

DOCUMENT TYPE: Patent LANGUAGE: Chinese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|-----------|----------|---------------------------|-----------|
| | | | | |
| CN 1210019 | A | 19990310 | CN 1998-117763 | 19980910 |
| PRIORITY APPLN. INFO.: | | | CN 1998-117763 | 19980910 |
| AB The title process | comprises | cleaning | bovine tendon, sterilizin | a, treati |

The title process comprises cleaning bovine tendon, sterilizing, treating with protease, then treating successively with acid, base and organic solvent to obtain pure collagen, crosslinking, and drying at (-40)-35°. The protease is selected from pepsase, papain, trypsin, and bromelin; the base from NaOH, KOH, NaHCO3, and Na2CO3; the acid from formic acid, acetic acid, malonic acid, and citric acid; the organic solvent from methanol, ethanol, Et ether, acetone, and butanol; and the crosslinking agent from formaldehyde, acetaldehyde, and glutaraldehyde. The collagen sponge is useful for wound healing and as hemostatic.

L58 ANSWER 13 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1993:61902 CAPLUS

DOCUMENT NUMBER: 118:61902

TITLE: Collagen manufacture from intestines, ruminant

stomachs, lungs, and udders Sjoelander, E.

INVENTOR(S): PATENT ASSIGNEE(S):

Collagen Casing Einar Sjoelander AB, Swed.

SOURCE: Swed., 10 pp. CODEN: SSXXAY DOCUMENT TYPE: Patent

LANGUAGE: Swedish

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PA: | TENT I | | | | KIN |) | | | | APP | LICAT | | | | | ATE | |
|-----|--------|------|-----|-----|-----|-----|------|------|-----|-----|-------|------|-----|-----|-----|------|-----|
| SE | 4677 | | | | В | - | | 0907 | | SE | 1991- | 999 | | | | 910 | |
| | 9100 | | | | A | | 1992 | 0907 | | | | | | | | | |
| | 4677 | | | | | | 1993 | | | | | | | | | | |
| SE | 9200 | 649 | | | Ā | | 1992 | 1006 | | SE | 1992- | 649 | | | 1 | 9920 | 304 |
| CA | 2107 | 680 | | | A1 | | 1992 | 1006 | | CA | 1992- | 2107 | 680 | | 1 | 9920 | 326 |
| WO | 9217 | 503 | | | A1 | | 1992 | 1015 | | WO | 1992- | SE19 | 2 | | 15 | 9920 | 326 |
| | W: | AT, | AU, | BB, | BG, | BR, | CA, | CH, | CS, | DE | , DK, | ES, | FI, | GB, | HU, | JP, | KP, |
| | | KR, | LK, | LU, | MG, | MN, | MW, | NL, | NO, | PL | , RO, | RU, | SD, | SE, | US | | |
| | RW: | AT, | BE, | BF, | ВJ, | CF, | CG, | CH, | CI, | CM | , DE, | DK, | ES, | FR, | GA, | GB, | GN, |
| | | GR, | IT, | LU, | MC, | ML, | MR, | NL, | SE, | SN | , TD, | TG | | | | | |
| AU | 9214 | 221 | | | A | | 1992 | 1102 | | ΑU | 1992- | 1422 | 1 | | 1 | 9920 | 326 |
| EP | 5786 | 61 | | | A1 | | 1994 | 0119 | | EΡ | 1992- | 9069 | 06 | | 1 | 9920 | 326 |
| EP | 5786 | 61 | | | В1 | | 1996 | 0911 | | | | | | | | | |
| | R: | AT, | BE, | CH, | DE, | DK, | ES, | FR, | GB, | GR | , IT, | LI, | LU, | NL | | | |
| JP | 0650 | 5982 | | | T | | 1994 | 0707 | | JΡ | 1992- | 5065 | 89 | | 1: | 9920 | 326 |
| | 1426 | | | | | | 1996 | 0915 | | AΤ | 1992- | 9069 | 06 | | 1 | 9920 | 326 |
| ES | 2094 | 904 | | | Т3 | | 1997 | 0201 | | ES | 1992- | 9069 | 06 | | 1 | 9920 | 326 |
| NO | 9303 | 507 | | | A | | 1993 | 0930 | | NO | 1993- | 3507 | | | 1: | 9930 | 930 |

| RU 2094999 | C1 | 19971110 | RU | 1993-58205 | | 19931004 |
|------------------------|----|----------|----|-------------|---|----------|
| US 5411887 | A | 19950502 | US | 1993-133083 | | 19931005 |
| PRIORITY APPLN. INFO.: | | | SE | 1991-999 | Α | 19910405 |
| | | | | 1992-SE192 | | 19920326 |
| | | | | | | |

AB The process comprises cleaning the starting material, immersing the material in ice water, adjusting the pH to 5.5, grinding the mixture of ice water and starting material, adding addnl. water in an amount such that the ground mixture contains approx. equal amts. of starting material and water, heating the mixture to 40-42° and adjusting the pH to ≤11, preferably 10.5, and adding a proteolytic enzyme in an amount corresponding to 60 Anson units/kg dry solids to allow the hydrolysis of proteins other than collagen, maintaining the pH by addition of alkali until the hydrolysis is completed, adjusting the pH to 5.5 by addition of acid, and separating and collecting the collagen. Clear, transparent films are obtained by mixing the collagen with a reducing agent, e.g., ascorbic acid or NaHSO3, 52, crosslinking agent, e.g., glutaraldehyde, .apprx.0.1, and plasticizer, i.e, glycerin, 5-10 weight% (all based on dry collagen).

L58 ANSWER 14 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1991:150178 CAPLUS

DOCUMENT NUMBER: 114:150178

TITLE: Manufacture of microcapsules from atelocollagen and polyholosides for cosmetic, pharmaceutical or food

compositions

INVENTOR(S): Levy, Marie Christine; Andry, Marie Christine; Huc,

Alain; Buffevant, Chantal

PATENT ASSIGNEE(S): Bioetica S. A., Fr.
SOURCE: Eur. Pat. Appl., 16 pp.

SOURCE: Eur. Pat. Appl., 16 pp. CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PAT | TENT NO. | | KIND | DATE | APPLICATION NO. | DATE |
|-----|----------|--------|------|----------|------------------------|------------|
| | 381543 | | A1 | 19900808 | EP 1990-400030 | 19900105 |
| EP | 381543 | | B1 | 19930526 | | |
| | R: AT, | BE, CH | | | GB, GR, IT, LI, LU, NL | |
| | 2642329 | | A1 | 19900803 | FR 1989-1221 | 19890131 |
| FR | 2642329 | | B1 | 19910524 | | |
| AT | 89766 | | T | 19930615 | AT 1990-400030 | 19900105 |
| ES | 2058827 | | Т3 | 19941101 | ES 1990-400030 | 19900105 |
| AU | 9048864 | | A | 19900809 | AU 1990-48864 | 19900129 |
| AU | 633866 | | B2 | 19930211 | | |
| CA | 2009065 | | A1 | 19900731 | CA 1990-2009065 | 19900131 |
| CA | 2009065 | | С | 19990824 | | |
| JP | 02229111 | | A | 19900911 | JP 1990-21927 | 19900131 |
| JP | 2534921 | | B2 | 19960918 | | |
| KR | 163171 | | B1 | 19981201 | KR 1990-1111 | 19900131 |
| US | 5395620 | | A | 19950307 | US 1993-74701 | 19930608 |
| | 5622656 | | A | 19970422 | US 1994-328903 | 19941025 |
| | APPLN. I | NFO. : | | | FR 1989-1221 A | |
| | | | | | US 1989-336711 A | |
| | | | | | EP 1990-400030 A | |
| | | | | | | 1 19910826 |
| | | | | | | 3 19930608 |

AB The microcapsules of the invention comprise a mixed wall of crosslinked atelocollagen and polyholosides (e.g. glycosaminoglycans), the proportion

of the latter relative to the atelocollagen being preferably 18-50 weight%. The microcapsules can be manufactured either by a process involving interfacial crosslinking or by extrusion of a laminar flow which is broken up by vibrations into individual droplets, which fall in a crosslinking bath. The atelocollagen-containing microcapsules are biocompatible and are especially suitable for the manufacture of cosmetic, pharmaceutical, or food compns. Manufacture of microcapsules containing vitamin C, CD RED 30 pigment, olive

oil, salmon oil, or oenethera oil is described.

L58 ANSWER 15 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1985:506605 CAPLUS
DOCUMENT NUMBER: 103:106605

DOCUMENT NUMBER: 103:106605 ORIGINAL REFERENCE NO.: 103:17081a,17084a

TITLE: Shaped product of collagen by syneresis

INVENTOR(S): Yoden, Yoshimasa; Okuda, Tsuneo; Fuchigami, Eiji;

Kuwabara, Toshihiro
PATENT ASSIGNEE(S): Nitta Gelatin Co., Ltd., Japan

SOURCE: Eur. Pat. Appl., 17 pp.
CODEN: EPXXDW

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | | KIND | DATE | APPLICATION NO. | DATE |
|-------------------------------------|--------|----------------|----------------------------------|--------------------------------|----------------------|
| EP 143512
EP 143512
EP 143512 | | A1
B1
B2 | 19850605
19880330
19910710 | EP 1984-305534 | 19840814 |
| | R, GB, | | 19850806 | US 1984-632855 | 19840720 |
| AU 8433461
AU 569112 | | A
B2 | 19850418
19880121 | AU 1984-33461 | 19840924 |
| ES 536242
FI 8403840 | | A1
A | 19850716
19850330 | ES 1984-536242
FI 1984-3840 | 19840926
19840928 |
| FI 77678
FI 77678 | | B
C | 19881230
19890410 | | |

PRIORITY APPLN. INFO.: JP 1983-182437 A 19830929
AB Shaped products are prepared from collagen by applying a crosslinking

agent to the pasty collagen composition being shaped, freezing the shaped product to enable the crosslinking reaction by separation of water, and thawing the crosslinked product. Thus, a fresh corium layer of unshayed oxhide was dipped for 10 days in 2 parts of 0.4% lime milk per 1 part corium, washed, neutralized by HCl. dipped 5 h in 2 parts 18 agueous NH4Cl solution

per 1

part corium, washed, and ground to give collagen fibers. An aqueous suspension containing the corium at 8% solids concentration and NaOH at 3% was repared

from 20% of the fibers and kept at 20° for 2 days. HCl was added to the emulsion at 520° to lower the pH to 4.0 and precipitate a fibrous agglomerate which was dehydrated. The remaining 80% of the fibers was added to the dehydrated product and swollen in aqueous citric acid at pH 3.0 at a solids concentration of 3.5%. The homogeneous mixture was

homogenized to form a pasty composition which was extruded through an annular nozzle into a 20% saline coagulating solution containing 1000 ppm glutaraldehyde

[111-30-8]at

pH 9.5 and 20 $^{\rm e}$. The extruded tube had pH 3.6, and it was left in the solution for 20 min. until its pH increased to 9.0, washed in flowing

water for 10 min, placed in a freezer at -20°, and kept frozen for 5 h. The tube had water content 96% before freezing and 75% after freezing and thawing. The bursting strength of the tube increased from 600 to 1500 mm H2O/cm2 after freezing and thawing. The wet tube was filled with sausage meat, dried at 75° for 20 min, and boiled 20 min at 80° to make a sausage which was cooked in a frying pan without breaking the tube.

L58 ANSWER 16 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1984:557716 CAPLUS DOCUMENT NUMBER: 101:157716 ORIGINAL REFERENCE NO.: 101:23799a,23802a

TITLE:

Collagen fleece INVENTOR(S):

Paques, Eric Paul; Fuhge, Peter PATENT ASSIGNEE(S): Behringwerke A.-G., Fed. Rep. Ger.

SOURCE: Ger. Offen., 11 pp. CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|--------|----------|-------------------|----------|
| | | | | |
| DE 3248188 | A1 | 19840628 | DE 1982-3248188 | 19821227 |
| EP 114351 | A2 | 19840801 | EP 1983-112861 | 19831221 |
| EP 114351 | A3 | 19850717 | | |
| EP 114351 | B1 | 19890726 | | |
| R: AT, CH, DE, | FR, IT | , LI | | |
| AT 44878 | T | 19890815 | AT 1983-112861 | 19831221 |
| JP 59133276 | A | 19840731 | JP 1983-244360 | 19831226 |
| ES 528400 | A1 | 19850116 | ES 1983-528400 | 19831226 |
| PRIORITY APPLN. INFO.: | | | DE 1982-3248188 A | 19821227 |
| | | | EP 1983-112861 A | 19831221 |

A collagen-containing material is treated with a neutral salt solution, a citric acid [77-92-9] solution, a solution of pepsin [9001-75-6], contacted with an ion exchanger, and the collagen is precipitated with a neutral salt, treated with a crosslinking agent, and dried to give a wound covering. Thus, 5 kg residue from the extraction of chopped placentas with isotonic saline was minced, extracted with pH 7.4 0.05 M Tris-HCl buffer containing 2 M NaCl, the residue washed with H2O at 4°, suspended in 1M citric acid for 1 h, and the residue was homogenized with H2O at 4°, centrifuged, suspended in 0.1M HOAc, and adjusted to pH 2 with HCl. The suspension was incubated twice with pepsin for 24 h at 25°, mixed with Dicalite, homogenized, and centrifuged. The supernatant was adjusted to pH 8 with Tris, stirred with Dowex 2-X8 for 1 h, solid NaCl was added to 0.2M, the mixture was centrifuged, the residue in H2O was brought to pH 5 with HOAc and dialyzed against H2O to give white collagen fibrils. The fibrils could be homogenized in H2O at pH 5, brought to pH 8, treated with H2CO [50-00-0] at 25°, and freeze-dried to give a fleece with good H2O absorption, strength, and elasticity.

L58 ANSWER 17 OF 17 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1984:460178 CAPLUS DOCUMENT NUMBER: 101:60178

ORIGINAL REFERENCE NO.: 101:9259a,9262a TITLE:

Collagen-glycosaminoglycan composite materials INVENTOR(S): Yannas, Ioannis V.; Kirk, James F. PATENT ASSIGNEE(S): Massachusetts Institute of Technology, USA

SOURCE: U.S., 8 pp.
CODEN: USXXAM
DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

US 4448718 A 19840515 US 1983-531804 19830913
PRIORITY APPLIN. INFO.: 1983-531804 19830913

AB A crosslinked collagen-glycosaminoglycan composite [Mc (average mol. weight of the segments between adjacent crosslinks) = 800-10,000] is prepared from contacting the uncrosslinked composite with a gaseous aldehyde. Artificial skin produced by this method is more stable toward long-term storage than similar materials produced by other methods. Thus, a

storage than similar materials produced by other methods. Thus, a collagen dispersion was prepared by contacting strips of calf hide with an aqueous solution containing propionic acid and benzoic acid. The collagen was purified by a precipitation process, then dispersed in a citric acid-buffer solution at pH 3.2. The dispersion was copptd. with a 1% chondroitin 6-sulfate solution (oH 3.2); the precipitate was homogenized, filtered, and

dried.

A composite material prepared from the above mixture in the form of a sheet was crosslinked with glutaraldehyde [111-30-8] vapor generated from a 25% glutaraldehyde solution in a vented desiccator. The treated sheets had a much lower Mc than untreated sheets.

=> d his

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(FILE 'HOME' ENTERED AT 09:04:35 ON 31 JAN 2008)

FILE 'REGISTRY' ENTERED AT 09:04:41 ON 31 JAN 2008

E CITRIC ACID E CITRIC ACID/CN

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E CITRIC ACID/CN

L2 15758 S E 3 E OXALACETIC ACID/CN

L3 1 S E3

E CITRIC ACID/CN

1 S E3 E ACONITIC ACID/CN

L5 1 S E3 E MALATE

5352 S E3

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L8 22764 S L1 L9 920942 S L2

L10 4146 S L3 L11 68175 S L4 L12 1003 S L5

L13 22725 S L6

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E N-HYDROXYSUCCINIMIDE/CN

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STN Search - 10/517,692

L14

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L16

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L19

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L21

L22

L23

1 S E3

1 S E3

5280 S L14

312 S L15

5501 S L16 OR L17

162 S L15 AND (PY<=2003)

3763 S L10 AND (PY<=2003)

784186 S L7 AND (PY<=2003)

18251 S L8 AND (PY<=2003)

707903 S L9 AND (PY<=2003)

E N-HYDROXYSULFOSUCCINIMIDE/CN

FILE 'CAPLUS' ENTERED AT 09:10:58 ON 31 JAN 2008

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STN Search - 10/517,692
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L55
              1 S L53 AND POLYCARBOXYLIC ACID
L56
             6 S L53 AND CARBOXYLIC ACID
           617 S L46 AND CITRIC ACID
L57
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         72141 GELATIN
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         83422 GELATIN
                (GELATIN OR GELATINS)
L61
             4 L59 AND GELATIN
=> d ibib abs 1-4
L61 ANSWER 1 OF 4 CAPLUS COPYRIGHT 2008 ACS on STN
ACCESSION NUMBER:
                        2007:384892 CAPLUS
DOCUMENT NUMBER:
                         146:374899
TITLE:
                         Immobilization of enzymes by adsorption on porous
                         carrier with subsequent crosslinking in the presence
                         of a polyfunctional amine for use in organic synthesis
INVENTOR(S):
                         Mazeaud, Isabelle; Poulsen, Poul Boerge Rosenius;
                         Christensen, Morten Wuertz; Brask, Jesper
PATENT ASSIGNEE(S):
                        Novozymes A/S, Den.
                         PCT Int. Appl., 32pp.
SOURCE:
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
     PATENT NO.
                        KIND DATE APPLICATION NO. DATE
                         A1 20070405 WO 2006-DK542 20061002
     WO 2007036235
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,
             CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
             GE, GH, GM, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LV, LY, MA, MD, MG, MK, MN,
             MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RS,
             RU, SC, SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN, TR, TT, TZ,
             UA, UG, UZ, VC, VN, ZA, ZM, ZW
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IS, IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ,
CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH,
GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
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KG, KZ, MD, RU, TJ, TM

US 2007087418 20070419 US 2006-541615 A1 20061002 PRIORITY APPLN. INFO .: DK 2005-1368 A 20050930 US 2005-724862P P 20051007

The present invention relates to the immobilization of enzymes by adsorbing enzymes, a polyfunctional amine and a crosslinking agent onto a particulate porous carrier in a mixer apparatus or in a fluid bed apparatus

The function of the polyfunctional amine is to provide a network of amine-groups available for covalent crosslinking with the crosslinking agent and the enzymes amine-groups. In particular, immobilization of lipase B on a silica-based carrier by impregnation and subsequent crosslinking by glutaraldehyde in the presence of polyethylene imine is described. The immobilized enzyme of the invention is useful for modification of organic compds. such as esterification, epoxidn., hydrolysis or ring opening.

REFERENCE COUNT: THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L61 ANSWER 2 OF 4 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2000:529535 CAPLUS

DOCUMENT NUMBER: 133:137057

TITLE: Hot water-resistant gelatin gels useful as

biomaterials

INVENTOR(S): Nagura, Masanobu; Mochizuki, Akira

PATENT ASSIGNEE(S): Terumo Corp., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 4 pp.

CODEN: JKXXAF DOCUMENT TYPE: Patent

LANGUAGE: Japanese FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|---------|-------------|------------------------|----------|
| | | | | |
| JP 2000212286 | A | 20000802 | JP 1999-16038 | 19990125 |
| PRIORITY APPLN. INFO.: | | | JP 1999-16038 | 19990125 |
| AB The gels are obtain | ed from | gelatin sub | stance by crosslinking | with |

polycarboxylic acids under heat and have swelling ratio (Sc) ≤ 1.0 where Sc = $\{Sc(40)/Sc(30)\}/\{Sh(40)/Sh(30)\}$ and 30 and 40 are water temperature in degree (swelling degree Sc and Sh are derived from S = ((Ws-Wd)/Wd; Ws = weight of gel at equilibrium state; Wd = weight of gel at dry state). Examples of polycarboxylic acids are succinic acid, citric acid, and adipic acid.

L61 ANSWER 3 OF 4 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1994:53389 CAPLUS

DOCUMENT NUMBER: 120:53389

Ionic complexes of ionizable emulsifiers with TITLE:

ionizable polypeptides and/or ionizable hydrocolloids INVENTOR(S): Reimer, Robert A.; Carruthers, Mark S.; Corr, Robert

J., Jr.; Miller, James W.; Tarlton, Eugene

PATENT ASSIGNEE(S):

Pfizer Inc., USA SOURCE: PCT Int. Appl., 37 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

| PA | TENT | NO. | | | KIN |) | DATE | | AP | PLICAT | ION | NO. | | Ε | ATE | |
|---------|-------|------|------|-----|-----|-----|------|------|-------|--------|------|-----|-----|------|------|-----|
| | | | | | | - | | | | | | | | - | | |
| WO | 9321 | 784 | | | A1 | | 1993 | 1111 | WO | 1993- | US21 | 67 | | 1 | 9930 | 316 |
| | W: | AU, | CA, | JP, | KR, | NO, | RU, | UA, | US | | | | | | | |
| | RW: | AT, | BE, | CH, | DE, | DK, | ES, | FR, | GB, G | R, IE, | IT, | LU, | MC, | NL, | PT, | SE |
| AU | 9339 | 169 | | | A | | 1993 | 1129 | AU | 1993- | 3916 | 9 | | 1 | 9930 | 316 |
| EP | 6372 | 09 | | | A1 | | 1995 | 0208 | EP | 1993- | 9082 | 96 | | 3 | 9930 | 316 |
| | R: | AT, | BE, | CH, | DE, | DK, | ES, | FR, | GB, G | R, IE, | IT, | LI, | LU, | NL, | PT, | SE |
| JP | 0750 | 2172 | | | T | | 1995 | 0309 | JP | 1993- | 5192 | 46 | | 1 | 9930 | 316 |
| IL | 1054 | 80 | | | A | | 1997 | 0110 | IL | 1993- | 1054 | 80 | | 1 | 9930 | 415 |
| ZA | 9302 | 839 | | | A | | 1994 | 1024 | ZA | 1993- | 2839 | | | 1 | 9930 | 422 |
| NO | 9404 | 012 | | | A | | 1994 | 1021 | NO | 1994- | 4012 | | | 1 | 9941 | 021 |
| PRIORIT | Y APP | LN. | INFO | . : | | | | | US | 1992- | 8728 | 69 | 2 | A1 1 | 9920 | 423 |
| | | | | | | | | | WO | 1993- | US21 | 67 | 7 | λ 1 | 9930 | 316 |

AB Complexes of ionizable emulsifiers with ionizable polypeptides and ionizable hydrocolloids are described for use as fat substitutes, food opacifiers, foam stabilizers and flavor modifiers. They are further useful as stiffeners for oils and oil-water emulsions allowing the use of normally liquid unsatd. oils in place of saturated fats in food compns. such as shortenings and spreads. Whey protein concentrate 40 was dissolved in water

600

g and a mixture of stearic acid 60% and palmitic acid 40% 100 g was added with stirring and heating to 75°. The pH of the mixture was adjusted to pH 6.8 with NaOH to form an opaque, viscous solution that after cooling and refrigeration had the appearance, odor, and texture of soft fat. The use of the fat substitutes of the invention in spreads, frosting, desserts, mayonnaise etc. is demonstrated.

L61 ANSWER 4 OF 4 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1988:210254 CAPLUS

DOCUMENT NUMBER: 108:210254

DOCUMENT NUMBER: 108:210254

TITLE: Process for manufacture of crosslinked gelatin-impregnated vascular grafts

INVENTOR(S): Maini, Roshan

PATENT ASSIGNEE(S): Vascutek Ltd., UK

SOURCE: Pat. Specif. (Aust.), 11 pp.

CODEN: ALXXAP
DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | KIND DATE | APPLICATION NO. | DATE |
|------------------------------------|--------------------------------|---------------------------------|------------------------|
| AU 569645
AU 8550593 | B2 19880211
A 19860605 | | 19851129 |
| EP 183365 | A3 19880406 | EP 1985-307255 | 19851010 |
| EP 183365
R: AT, BE, CH, | B1 19921230
DE, FR, GB, IT, | LI, NL, SE | |
| AT 83911 | T 19930115 | | 19851010 |
| DK 172304 | B1 19980309 | | 19851128 |
| JP 02011258 PRIORITY APPLN. INFO.: | В 19900313 | JP 1985-267619
GB 1984-30265 | 19851129
A 19841130 |
| TRIUNIII MEPLN. INFU.: | | EP 1985-307255 | A 19851010 |

AB Vascular grafts, which require no blood preimpregnation, and which after implantation start to degrade and become permeable at a known rate so that tissue ingrowth can take place, are prepared by impregnating a tube of a

L24

porous flexible material with gelatin and treating the impregnated tube with an amino group crosslinking agent. A tube formed as a knitted textile material structure was impregnated under vacuum with a mixture of a gelatin which had been treated with succinoyl chloride to cause crosslinking of 75% of its free amino groups and untreated gelatin (mole ratio 1:1) at 65°. The gelatin mixture was allowed to gel, and tubes subjected to a treatment with a 20% HCHO solution at pH 4 and 4° for 16 h, and the formed vascular graft washed 5 times in pyrogen-free H2O at room temperature. This graft became fully porous after 25-30 h under laboratory

test conditions. Comparison grafts prepared using untreated gelatin only became fully porous in $>45\ h.$

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50287 S L11 AND (PY<=2003)

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            48 S L40 AND L41
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L47
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L48
           2066 S L46 AND L47
L49
              8 S CHRONDROITIN SULFATE
               E CHONDROITIN SULFATE+ALL/CT
L50
         13474 S (CHONDROITIN SULFATE OR "CHONDROITIN, HYDROGEN SULFATE")
           116 S L46 AND L50
L51
1.52
             8 S L51 AND BIOMATERIAL
    FILE 'CAPLUS' ENTERED AT 11:24:22 ON 31 JAN 2008
L53
           168 S L46 AND BIOMATERIAL
L54
             2 S L53 AND CITRIC ACID
L55
             1 S L53 AND POLYCARBOXYLIC ACID
             6 S L53 AND CARBOXYLIC ACID
L57
           617 S L46 AND CITRIC ACID
            17 S L57 AND COLLAGEN
L58
L59
           733 S L46 AND POLYCARBOXYLIC ACID
L60
            0 S L59 AND COLLAGEN
L61
             4 S L59 AND GELATIN
=> log h
COST IN U.S. DOLLARS
                                                SINCE FILE
                                                              TOTAL
                                                     ENTRY SESSION
FULL ESTIMATED COST
                                                     130.42
                                                              452.78
```

L13

22725 S L6

```
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)
                                              SINCE FILE
                                                              TOTAL.
                                                     ENTRY
                                                             SESSION
CA SUBSCRIBER PRICE
                                                     -24.00
                                                               -75.20
SESSION WILL BE HELD FOR 120 MINUTES
STN INTERNATIONAL SESSION SUSPENDED AT 11:42:47 ON 31 JAN 2008
Connecting via Winsock to STN
Welcome to STN International! Enter x:x
LOGINID: SSPTASYG1600
PASSWORD:
* * * * * * RECONNECTED TO STN INTERNATIONAL * * * * *
SESSION RESUMED IN FILE 'CAPLUS' AT 13:29:08 ON 31 JAN 2008
FILE 'CAPLUS' ENTERED AT 13:29:08 ON 31 JAN 2008
COPYRIGHT (C) 2008 AMERICAN CHEMICAL SOCIETY (ACS)
COST IN U.S. DOLLARS
                                                SINCE FILE
                                                                TOTAL
                                                     ENTRY
                                                            SESSION
FULL ESTIMATED COST
                                                    130.42
                                                               452.78
                                                               TOTAL
DISCOUNT AMOUNTS (FOR OUALIFYING ACCOUNTS)
                                               SINCE FILE
                                                    ENTRY
                                                             SESSION
CA SUBSCRIBER PRICE
                                                     -24.00
                                                                -75.20
=> d his
     (FILE 'HOME' ENTERED AT 09:04:35 ON 31 JAN 2008)
    FILE 'REGISTRY' ENTERED AT 09:04:41 ON 31 JAN 2008
               E CITRIC ACID
               E CITRIC ACID/CN
               E MALIC ACID/CN
L1
              1 S E3
               E CITRIC ACID/CN
L2
         15758 S E 3
               E OXALACETIC ACID/CN
1.3
              1 S E3
               E CITRIC ACID/CN
             1 S E3
L4
               E ACONITIC ACID/CN
              1 S E3
               E MALATE
L6
          5352 S E3
    FILE 'CAPLUS' ENTERED AT 09:09:21 ON 31 JAN 2008
L7
        1016908 S L1 OR L2 OR L3 OR L4 OR L5 OR L6
L8
         22764 S L1
1.9
        920942 S L2
T-10
         4146 S L3
         68175 S L4
1.12
         1003 S L5
```

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FILE 'REGISTRY' ENTERED AT 09:10:08 ON 31 JAN 2008
                E HYDROXYSUCCINIMIDE
               E N-HYDROXYSUCCINIMIDE/CN
L14
              1 S E3
              E N-HYDROXYSULFOSUCCINIMIDE/CN
              1 S E3
    FILE 'CAPLUS' ENTERED AT 09:10:58 ON 31 JAN 2008
L16
          5280 S L14
L17
           312 S L15
L18
          5501 S L16 OR L17
L19
           162 S L15 AND (PY<=2003)
L20
        784186 S L7 AND (PY<=2003)
L21
         18251 S L8 AND (PY<=2003)
L22
        707903 S L9 AND (PY<=2003)
L23
          3763 S L10 AND (PY<=2003)
L24
         50287 S L11 AND (PY<=2003)
L25
           890 S L12 AND (PY<=2003)
L26
          19656 S L13 AND (PY<=2003)
L27
              0 S L19 AND L25
L28
              8 S L19 AND L20
     FILE 'REGISTRY' ENTERED AT 09:17:53 ON 31 JAN 2008
               E MALIC ACID/CN
1.29
              0 S E3/RACT
    FILE 'CAPLUS' ENTERED AT 09:19:17 ON 31 JAN 2008
L30
          1540 S L1/RACT
L31
          46553 S L2/RACT
L32
           633 S L3/RACT
          4190 S L4/RACT
L33
L34
            42 S L5/RACT
L35
           800 S L6/RACT
L36
          4322 S L14/RACT
L37
           184 S L15/RACT
L38
          53062 S L30 OR L31 OR L32 OR L33 OR L34 OR L35
L39
          4454 S L36 OR L37
L40
          41864 S L38 AND (PY<=2003)
L41
          3152 S L39 AND (PY<=2003)
L42
             48 S L40 AND L41
     FILE 'CAPLUS' ENTERED AT 09:48:36 ON 31 JAN 2008
L43
                STRUCTURE UPLOADED
                S L43
    FILE 'REGISTRY' ENTERED AT 09:49:10 ON 31 JAN 2008
L44
             0 S L43
     FILE 'CAPLUS' ENTERED AT 09:49:10 ON 31 JAN 2008
L45
             0 S L44
     FILE 'CAPLUS' ENTERED AT 11:02:13 ON 31 JAN 2008
                E CROSSLINKING+ALL/CT
L46
         72274 S CROSSLINKING AGENT
1.47
         595653 S SULFATE
T.48
           2066 S L46 AND L47
L49
              8 S CHRONDROITIN SULFATE
               E CHONDROITIN SULFATE+ALL/CT
1.50
         13474 S (CHONDROITIN SULFATE OR "CHONDROITIN, HYDROGEN SULFATE")
L51
           116 S L46 AND L50
```

L52 8 S L51 AND BIOMATERIAL

FILE 'CAPLUS' ENTERED AT 11:24:22 ON 31 JAN 2008 168 S L46 AND BIOMATERIAL L54 2 S L53 AND CITRIC ACID L55 1 S L53 AND POLYCARBOXYLIC ACID L56 6 S L53 AND CARBOXYLIC ACID L57 617 S L46 AND CITRIC ACID L58 17 S L57 AND COLLAGEN L59 733 S L46 AND POLYCARBOXYLIC ACID L60 0 S L59 AND COLLAGEN

4 S L59 AND GELATIN

=> fil caplus

L61

 COST IN U.S. DOLLARS
 SINCE FILE ENTRY
 TOTAL SESSION SESSION 130.90

 FULL ESTIMATED COST
 130.90
 453.26

 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)
 SINCE FILE TOTAL SESSION 150.00
 TOTAL 150.00

 CA SUBSCRIBER PRICE
 -24.00
 -75.20

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FILE COVERS 1907 - 31 Jan 2008 VOL 148 ISS 5 FILE LAST UPDATED: 30 Jan 2008 (20080130/ED)

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http://www.cas.org/infopolicv.html

=> s 146 and polyacid 1639 POLYACID 988 POLYACIDS 2317 POLYACID (POLYACID OR POLYACIDS) L62 81 L46 AND POLYACID

=> s 162 and (py<+2003) 22927806 PY<+2003 L63 67 L62 AND (PY<+2003)

=> s 162 and (py<=2003)

23975525 PY<=2003 L64 68 L62 AND (PY<=2003) => s 164 and sulfate 546575 SULFATE

> 99691 SULFATES 595653 SULFATE

(SULFATE OR SULFATES)

L65 2 L64 AND SULFATE

=> d ibib abs 1-2

L65 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2001:900492 CAPLUS

DOCUMENT NUMBER: 136:38944

TITLE: Chipping- and corrosion-resistant and sound-insulating coating compositions containing coating components

recovered from coating booth water for automotive bodies

INVENTOR(S): Tanaka, Yoshito; Taniquchi, Hitoshi; Kurabayashi,

Osamu

PATENT ASSIGNEE(S): Nippon Oil and Fats Basf Coating K. K., Japan; Fuji

Heavy Industries Ltd.

SOURCE: Jpn. Kokai Tokkyo Koho, 9 pp.

CODEN: JKXXAF
DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

JP 2001342436 A 20011214 JP 2000-166399 20000602 <-JP 3910342 B2 20070425

PRIORITY APPLN. INFO.: P2000-166399 20000602

B The composition comprises (A) a thermoplastic resin, (B) a coating recovered from circulating water of coating booth, and (C) a water soluble resin. Thus, 28 parts Poly bd-R 45HT (butadiene rubber) was mixed with a polyester-based coating recycled from circulating water of coating booth 6.9, adipic acid-1,4-butamediol-hexadecenylsuccinic anhydride-isophthalic acid-trimellitic anhydride-trimethylolpropane copolymer dimethylethanolamine salt 11.3, PW 380 (mineral oil-type plasticizing agent) 7.7, Hakuenka CCR (calcium carbonate) 36.8, Barite BA (barium sulfate) 11, Duranate TPA-B 80E (blocked isocyanate) 7 and butyl Cellosolve 2 parts, applied to a precoated steel plate and cured at 140° for 30 min, showing good water, chipping and corrosion resistance and sound insulation.

L65 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1988:619521 CAPLUS

DOCUMENT NUMBER: 109:219521

TITLE: Photographic support material with antistatic

back-coating

INVENTOR(S): Saeverin, Eckehard; Tyrakowski, Hans Udo
PATENT ASSIGNEE(S): Schoeller, Felix, Jr., G.m.b.H. und Co. K.-G., Fed.

PATENT ASSIGNEE(S): Schoeller, Fe. Rep. Ger.

SOURCE: Ger. Offen., 5 pp.

DOCUMENT TYPE: Ger. OTIEN., 5 PR
CODEN: GWXXBX
Patent

LANGUAGE: FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | | DATE |
|--|-------------|-----------------------------------|---|--------------|----------------------------------|
| DE 3700183
EP 274017 | A1
A2 | 19880714
19880713 | DE 1987-3700183
EP 1987-116068 | | 19870106 <
19871031 < |
| EP 274017
EP 274017 | A3
B1 | 19900228
19920729 | | | |
| R: AT, BE, CH,
AT 78938
ES 2033285 | DE, ES
T | , FR, GB,
19920815
19930316 | GR, IT, LI, LU, NL,
AT 1987-116068
ES 1987-116068 | SE | 19871031 <
19871031 < |
| JP 63173044
US 5104779 | A
A | 19880716
19920414 | JP 1988-165
US 1989-380212 | | 19880105 <
19890714 < |
| PRIORITY APPLN. INFO.: | | | DE 1987-3700183
EP 1987-116068
US 1988-141925 | A
A
B2 | 19870106
19871031
19880106 |

- AB An antistatic photog, support showing low staining during transport through roller-transport development apparatus, a high abrasion resistance and stability in alkaline developer solns, good printability with com. printing inks, good writability, and good adhesive tape adhesion contains a backing layer from a composition containing: (1) a colloidal Al-modified silicic acid; (2)
- an alkali salt of an organic polyacid; (3) an aqueous dispersion of an alkyl acrylate copolymer having free carboxyl groups 1-10 mol% and free OR groups 0-20 mol%; and (4) a trifunctional aziridine as a crosslinking agent. The method of preparing the support comprises adding the components in a specific sequence and forming a layer on the backside of a support with the mixture Thus, a typical backlayer composition contained Ludox AM (colloidal Al-modified silicic acid), Bu acrylate-methacrylic acid-styrene copolymer, a trifunctional aziridine, and Na cellulose sulfate.

=> s 164 and biomaterial 10076 BIOMATERIAL

10076 BIOMATERIAL 10856 BIOMATERIALS

16264 BIOMATERIAL

(BIOMATERIAL OR BIOMATERIALS)

L66 0 L64 AND BIOMATERIAL

=> s 164 and gelatin

CA SUBSCRIBER PRICE

72141 GELATIN 30834 GELATINS

83422 GELATIN

(GELATIN OR GELATINS)

L67 0 L64 AND GELATIN

=> log h
COST IN U.S. DOLLARS
SINCE FILE TOTAL
ENTRY
FULL ESTIMATED COST
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)
ENTRY
ENTRY
ENTRY
SESSION
174.68

-76.80

-1.60

SESSION WILL BE HELD FOR 120 MINUTES STN INTERNATIONAL SESSION SUSPENDED AT 13:33:40 ON 31 JAN 2008